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(54) Title: SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES ENCODED THEREBY

#### (57) Abstract

The present invention provides DNA molecules that constitute portions of the genome of a plant, and polypeptides encoded thereby. The DNA molecules are useful for expressing a gene product, either as a promoter or as a structural gene or as an UTR or as a 3' termination sequence and are also useful in controlling expression of the target gene or as tools for genetic mapping or identification of a particular individual plant or for clustering of a group of plants with a common trait.

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# SEQUENCE-DETERMINED DNA FRAGMENTS AND CORRESPONDING POLYPEPTIDES ENCODED THEREBY

#### FIELD OF THE INVENTION

The present invention relates to isolated polynucleotides that encode all, or a portion of, a gene that is expressed and the corresponding polypeptide. The present invention also relates to isolated polynucleotides that encode regulatory regions of genes.

#### 10 DESCRIPTION OF THE RELATED ART

Efforts to map and sequence the genome of a number of organisms are in progress; a few complete genome sequences, for example those of *E. coli* and *Saccharomyces cerevisiae* are known (Blattner et al., *Science* 277:1453 (1997); Goffeau et al., *Science* 274:546 (1996)). The complete genome of a multicellular organism, *C. elegans*, has also been sequenced (See, the *C. elegans* Sequencing Consortium, *Science* 282:2012 (1998)). To date, no complete genome of a plant has been sequenced, nor has a complete cDNA complement of any plant been sequenced.

#### SUMMARY OF THE INVENTION

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The present invention comprises polynucleotides, such as complete cDNA sequences and/or sequences of genomic encompassing complete genes, portions of qenes, intergenic regions, hereinafter collectively referred to as "Sequence-Determined DNA Fragments" (SDFs), from particularly corn and Arabidopsis thaliana and polypeptides derived therefrom. In some instances, the SDFs span the entirety of a protein-coding segment. In some instances, the entirety of an mRNA is represented. Other objects of the invention are the control sequences, such as but not limited to promoters, that are also represented by SDFs of the invention. Complements of any sequence of the invention are also considered part of the invention.

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Other objects of the invention are polynucleotides comprising exon sequences, polynucleotides comprising intron sequences, polynucleotides comprising introns together with intron/exon junction sequences, 5**′** untranslated sequences, and 3' untranslated sequences of the SDFs of the present invention. Polynucleotides representing the joinder of any exons described herein, in any arrangement, for example, to produce a sequence encoding any desirable amino acid sequence are within the scope of the invention.

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The present invention also resides in probes useful for isolating and identifying nucleic acids that hybridize to an SDF of the invention. The probes are typically of a length of 12 to 2000 nucleotides long; more typically, 15 to 200 nucleotides long; even more typically, 18 to 100 nucleotides long.

Yet another object of the invention is a method of isolating and/or identifying nucleic acids using the following steps:

- (a) contacting a probe of the instant invention with a polynucleotide sample under conditions that permit hybridization and formation of a polynucleotide duplex; and
  - (b) detecting and/or isolating the duplex of step (a).

The conditions for hybridization can be from low to moderate to high stringency conditions. The sample can include a polynucleotide having a sequence unique in a plant genome. Probes and methods of the invention are useful, for example, without limitation, for mapping of genetic traits and/or for positional cloning of a desired portion of genomic DNA.

Probes and methods of the invention can also be used for detecting alternatively spliced messages within a species. Probes and methods of the invention can further be used to detect or isolate related genes in other plant species using genomic DNA (gDNA) and/or cDNA libraries. In some instances, especially when longer probes and low to moderate stringency

hybridization conditions are used, the probe will hybridize to a plurality of cDNA and/or gDNA sequences of a plant. This approach is useful for isolating representatives of gene families which are identifiable by possession of a common functional domain in the gene product or which have common cisacting regulatory sequences. This approach is also useful for identifying orthologous genes from other organisms, which can be more or less related to corn, Arabidopsis, or another plant.

The present invention also resides in constructs for modulating the expression of the genes comprised of all or a portion of an SDF. The constructs comprise all or a portion of the expressed SDF, or of a complementary sequence. Examples of constructs include ribozymes comprising RNA encoded by an SDF or by a sequence complementary thereto, antisense constructs, constructs comprising coding regions or parts thereof, constructs comprising promoters, introns, untranslated regions, etc. When inserted into a host cell the construct is, preferably, functionally integrated with or operatively linked to a heterologous polynucleotide. For instance, a coding region from an SDF might be operably linked to a promoter that is functional in a plant.

The present invention also resides in host cells, including bacterial or yeast cells or plant cells, and transgenic plants that harbor constructs such as described above. Another aspect of the invention relates to methods for modulating expression of specific genes in transgenic plants by expression of the structural gene component of the constructs, by regulation of expression of one or more endogenous genes in a transgenic plant or by suppression of expression of the polynucleotides of the invention in a transgenic plant. Methods of modulation of gene expression include without limitation (1) inserting into a host cell additional copies of a polynucleotide comprising a coding sequence; (2) modulating an endogenous promoter in a host cell; (3) inserting antisense

or ribozyme constructs into a host cell and (4) inserting into a host cell a polynucleotide comprising a sequence encoding a mutant, fragment, or fusion of the native polypeptides of the instant invention.

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#### BRIEF DESCRIPTION OF THE TABLES

The sequences of exemplary SDFs and polypeptides encoded thereby of the instant invention are listed in SEQ TABLES 1 and 2; annotation relevant to the sequences shown in SEQ TABLES 1 and 2 is presented in REF TABLES 1 and 2. Each sequence corresponds to a Maximum Length cDNA Polynucleotide Sequence. Each Maximum Length cDNA Polynucleotide Sequence (Ac) corresponds to at least one sequence in SEQ TABLE 1 and 2. REF TABLE 1 corresponds with SEQ TABLE 1; REF TABLE 2 corresponds with SEQ TABLE 2.

REF TABLES 1 and 2 are Reference Tables which correlate each of the sequences and SEQ ID NOS in SEQ TABLES 1 and 2 with a corresponding Maximum Length cDNA Sequence (Ac), Ceres (Applicant) sequence identifier and other information about the individual sequence. SEQ TABLES 1 and 2 are Sequence Tables containing the sequence of each nucleic acid and amino acid sequence.

1 and 2, TABLES each section begins identifying the Maximum Length cDNA Polynucleotide Sequence, "Clone ID" that is а number used identification purposes by the applicant and in instances a "Public Genomic DNA" sequence, indicated by a "gi In those instances where a public sequence is recited, there follows information about gene annotations predicted exons. In this portion, after the description of functional unit, the starting gene's and nucleotide number of the public sequence and the computer program used to generate the result are listed. "INIT" denotes an initial exon. "INTR" denotes an internal exon.

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"TERM" denotes a terminal exon. For those sequences where the computer program is identified as "OCKHAM", "INIT" denotes the first detected exon and "TERM" denotes the last detected exon. "SINGLE" denotes the entire mRNA coding portion is in one exon. Other notations are considered intuitive to the skilled artisan.

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In the next section (Ac), the cDNA MLS is identified by its SEQ ID NO ("Pat. Appln. SEQ ID NO") and the Ceres sequence identifier ("Ceres seq id"), which is also merely an identifier useful for the applicant. The designation of "Alternative transcription start sites" can include both positive and negative numbers. Positive numbers refer to the referenced SEO ID NO directly. The positions indicated by negative numbers, if any, refer to positions in the public genomic sequences. In instances where there is a "Public Genomic DNA" referenced, the relevant genomic sequence can be by direct reference to the nucleotide indicated by the "gi" number shown for the Public Genomic DNA. In instances where there is no Public Genomic DNA referenced, the relevant nucleotide sequence for alignment is nucleotide sequence associated with the amino acid the sequence designated by a "gi" number in the section (Dp). these instances, the nucleotide sequence is found in GENBANK by clicking on the link in the National Center Biotechnology Information Entrez database. The numbering is relative to position 1 as determined by aligning the first residue of the MLS cDNA sequence (SEQ ID NO \*) with the genomic sequence corresponding to the relevant "gi" number.

Subsection (B) lists SEQ ID NOS and Ceres seq\_ids for polypeptide sequences encoded by the cDNA sequence and the location of the start codon within the cDNA sequence that codes for the polypeptide. Subsection (B) also describes additional features within the polypeptide sequence.

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Subsection (C) provides information regarding identified domains (where present) within the polypeptide and (where present) a name for the polypeptide. Subsection (Dp) provides (where present) information concerning amino acid sequences 5 that are found to be related and have some sequence identity to the polypeptide sequences of SEQ TABLES 1 and 2. "related" sequences are identified by a "gi" number and are amino acid sequences in the publicly accessible BLAST onthe NCBI FTP web site (accessible databases 10 The database at the NCBI FTP site ncbi.nlm.gov/blast). utilizes the "gi" identifiers to assign by NCBI a unique identifier for each sequence in the databases, thereby providing a non-redundant database for sequences from various databases, including GenBank, EMBL, DBBJ (DNA Database of 15 Japan) and PDB (Brookhaven Protein Data Bank). Subsection (Ba), when present, describes a sequence as being considered plant-specific (i.e. a gene found only in a plant) or describes a bichemical activity for the protein encoded by the exemplary SDF. Subsection (Dn) provides polynucleotide 20 sequences (where present) related to the Maximum Length cDNA sequence.

#### DETAILED DESCRIPTION OF THE INVENTION

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The invention relates to (I) polynucleotides and methods of use thereof, such as

- IA. Probes, Primers and Substrates;
  - IB. Methods of Detection and Isolation;
    - B.1. Hybridization;
    - B.2. Methods of Mapping;
    - B.3. Southern Blotting;
    - B.4. Isolating cDNA from Related Organisms;
    - B.5. Isolating and/or Identifying Orthologous Genes
  - IC. Methods of Inhibiting Gene Expression
    - C.1. Antisense

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- C.2. Ribozyme Constructs;
- C.3. Co-Suppression;
- C.4. Other Methods to Inhibit Gene Expression
- ID. Methods of Functional Analysis;
- IE. Promoter Sequences and Their Use;
  - IF. UTR's and/or Intron Sequences and Their Use; and
  - IG. Coding Sequences and Their Use.

The specification also discloses (II) polypeptides including, without limitation, native proteins, mutants, fragments, and fusions. Antibodies to said polypeptides are also disclosed.

The specification also discloses (III) methods of modulating polypeptide production or activity. Examples of such methods include (i) suppressed, (ii) enhanced, and (iii) directed expression.

The specification also discloses (IV) gene constructs and construction of expression vectors, including (IVA) coding sequences and (IVB) promoters, and (IVC) Signal Peptides, (V) transformation procedures to illustrate the invention by way of examples.

#### I. Polynucleotides

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A number of the nucleotide sequences disclosed in SEQ TABLES 1 AND 2 herein as representative of the SDFs of the invention can be obtained by sequencing genomic DNA (gDNA) and/or cDNA from corn plants grown from HYBRID SEED # 35A19, purchased from Pioneer Hi-Bred International, Inc., Supply Management, P.O. Box 256, Johnston, Iowa 50131-0256.

Exemplified SDFs of the invention represent portions of the genome of corn or *Arabidopsis* and/or represent mRNA expressed from that genome. The isolated nucleic acid of the invention also encompasses corresponding portions of the genome

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and/or cDNA complement of other organisms as described in detail below.

Starting material for cDNA synthesis for the exemplary corn cDNA clones having sequences presented in SEQ TABLES 1 AND 2 was poly(A)-containing polysomal mRNAs from inflorescences and root tissues of corn plants grown from HYBRID SEED # 35A19. Male inflorescences and female (pre-and post-fertilization) inflorescences were isolated at various stages of development. Selection for poly(A) containing polysomal RNA was done using oligo d(T) cellulose columns, as described by Cox and Goldberg, "Plant Molecular Biology: A Practical Approach", pp. 1-35, Shaw ed., c. 1988 by IRL, Oxford.

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Tissues were or each organ was individually pulverized and frozen in liquid nitrogen. Next, the samples homogenized in the presence of detergents and centrifuged. The debris and nuclei were removed from the sample and more detergents were added to the sample. sample was centrifuged and the debris was removed. Then the sample was applied to a 2M sucrose cushion to isolate polysomes. The RNA was isolated by treatment with detergents followed by ethanol precipitation and proteinase K centrifugation. The polysomal RNA from the different tissues was pooled according to the following mass ratios: 15/15/1for male inflorescences, female inflorescences and root, respectively. The pooled material was then used for cDNA synthesis by the methods described below.

The quality and the integrity of the polyA+ RNAs were evaluated.

A number of the nucleotide sequences disclosed in SEQ TABLES 1 AND 2 herein as representative of the SDFs of the invention can also be obtained by sequencing genomic DNA from Arabidopsis thaliana, Wassilewskija ecotype or by sequencing cDNA obtained from mRNA from such plants as described below.

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This is a true breeding strain. Seeds of the plant are available from the Arabidopsis Biological Resource Center at the Ohio State University, under the accession number CS2360. Seeds of this plant were deposited under the terms and conditions of the Budapest Treaty at the American Type Culture Collection, Manassas, VA on August 31, 1999, and were assigned ATCC No. PTA-595.

Starting material for cDNA synthesis for the exemplary Arabidopsis cDNA clones having sequences presented in SEQ TABLES 1 AND 2 was polysomal RNA isolated from the top-most inflorescence tissues and roots of Arabidopsis Landsberg erecta (L. er.) also obtained from the Arabidopsis Biological Resource Center. Nine parts inflorescence to every part root was used, as measured by mass. pulverized and exposed to liquid nitrogen. Next, the sample homogenized in the presence of detergents and then centrifuged. The debris and nuclei were removed from the sample and more detergents were added to the sample. sample was centrifuged and the debris was removed and the sample was applied to a 2M sucrose cushion to isolate polysomal RNA. Cox et al., "Plant Molecular Biology: A Practical Approach", pp. 1-35, Shaw ed., c. 1988 by IRL, The polysomal RNA was used for cDNA synthesis by the methods described below. Polysomal mRNA was then isolated as described above for corn cDNA. The quality of the RNA was assessed electrophoretically.

Following preparation of the mRNAs from various tissues as described above, selection of mRNA with intact 5' ends and specific attachment of an oligonucleotide tag to the 5' end of such mRNA was performed using either a chemical or enzymatic approach. Both techniques take advantage of the presence of the "cap" structure, which characterizes the 5' end of most intact mRNAs and which comprises a guanosine generally methylated once, at the 7 position.

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The chemical modification approach involves the optional elimination of the 2', 3'-cis diol of the 3' terminal ribose, the oxidation of the 2', 3'-cis diol of the ribose linked to the cap of the 5' ends of the mRNAs into a dialdehyde, and the coupling of the such obtained dialdehyde to a derivatized oligonucleotide tag. Further detail regarding the chemical approaches for obtaining mRNAs having intact 5' ends are disclosed in International Application No. WO96/34981 published November 7, 1996.

The enzymatic approach for ligating the oligonucleotide tag to the intact 5' ends of mRNAs involves the removal of the phosphate groups present on the 5' ends of uncapped incomplete mRNAs, the subsequent decapping of mRNAs having intact 5' ends and the ligation of the phosphate present at the 5' end of the decapped mRNA to an oligonucleotide tag. Further detail regarding the enzymatic approaches for obtaining mRNAs having intact 5' ends are disclosed in Dumas Milne Edwards J.B. (Doctoral Thesis of Paris VI University, Le clonage des ADNc complets: difficultes et perspectives nouvelles. Apports pour l'etude de la regulation de l'expression de la tryptophane hydroxylase de rat, 20 Dec. 1993), EPO 625572 and Kato et al., Gene 150:243-250 (1994).

In both the chemical <del>or</del>and the enzymatic approach, the oligonucleotide tag has a restriction enzyme site (e.g. an EcoRI site) therein to facilitate later cloning procedures. Following attachment of the oligonucleotide tag to the mRNA, the integrity of the mRNA is examined by performing a Northern blot using a probe complementary to the oligonucleotide tag.

For the mRNAs joined to oligonucleotide tags using either the chemical or the enzymatic method, first strand cDNA synthesis is performed using an oligo-dT primer with reverse transcriptase. This oligo-dT primer can contain an internal tag of at least 4 nucleotides, which can be different from one mRNA preparation to another. Methylated dCTP is used for cDNA

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first strand synthesis to protect the internal EcoRI sites from digestion during subsequent steps. The first strand cDNA is precipitated using isopropanol after removal of RNA by alkaline hydrolysis to eliminate residual primers.

Second strand cDNA synthesis is conducted using a DNA polymerase, such as Klenow fragment and a primer corresponding to the 5' end of the ligated oligonucleotide. The primer is typically 20-25 bases in length. Methylated dCTP is used for second strand synthesis in order to protect internal EcoRI sites in the cDNA from digestion during the cloning process.

Following second strand synthesis, the full-length cDNAs are cloned into a phagemid vector, such as pBlueScript<sup>TM</sup> (Stratagene). The ends of the full-length cDNAs are blunted with T4 DNA polymerase (Biolabs) and the cDNA is digested with EcoRI. Since methylated dCTP is used during cDNA synthesis, the EcoRI site present in the tag is the only hemi-methylated site; hence the only site susceptible to EcoRI digestion. In some instances, to facilitate subcloning, an Hind III adapter is added to the 3' end of full-length cDNAs.

The full-length cDNAs are then size fractionated using either exclusion chromatography (AcA, Biosepra) or electrophoretic separation which yields 3 to 6 different fractions. The full-length cDNAs are then directionally cloned either into pBlueScript™ using either the EcoRI and SmaI restriction sites or, when the Hind III adapter is present in the full-length cDNAs, the EcoRI and Hind III restriction sites. The ligation mixture is transformed, preferably by electroporation, into bacteria, which are then propagated under appropriate antibiotic selection.

Clones containing the oligonucleotide tag attached to full-length cDNAs are selected as follows.

The plasmid cDNA libraries made as described above are purified (e.g. by a column available from Qiagen). A positive selection of the tagged clones is performed as follows.

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Briefly, in this selection procedure, the plasmid DNA is converted to single stranded DNA using phage F1 gene ΙI endonuclease in combination with an exonuclease (Chang et al., Gene 127:95 (1993)) such as exonuclease III or T7 gene 6 exonuclease. The resulting single stranded DNA is purified using paramagnetic beads as described by Fry et al., Biotechniques 13: 124 (1992). Here the single stranded DNA is hybridized with a biotinylated oligonucleotide having a sequence corresponding to the 3' end of the oligonucleotide Preferably, the primer has a length of 20-25 bases. Clones including a sequence complementary to the biotinylated oligonucleotide are selected by incubation with streptavidin coated magnetic beads followed by magnetic capture. capture of the positive clones, the plasmid DNA is released from the magnetic beads and converted into double stranded DNA using a DNA polymerase such as ThermoSequenase™ (obtained from Amersham Pharmacia Biotech). Alternatively, protocols such as the Gene Trapper<sup>TM</sup> kit (Gibco BRL) can be used. The double stranded DNA is then transformed, preferably electroporation, into bacteria. The percentage of positive clones having the 5' tag oligonucleotide is typically estimated to be between 90 and 98% from dot blot analysis.

Following transformation, the libraries are ordered in microtiter plates and sequenced. The Arabidopsis library was deposited at the American Type Culture Collection on January 7, 2000 as "E-coli liba 010600" under the accession number \_\_\_\_\_\_\_

Other methods for cloning full-length CDNA described, for example, by Seki et al., Plant Journal 15:707-(1998) "High-efficiency cloning of Arabidopsis fulllength cDNA by biotinylated Cap trapper"; Maruyama et al., Gene 138:171 (1994) "Oligo-capping a simple method to replace the cap structure eukaryotic of mRNAs with oligoribonucleotides"; and WO 96/34981.

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It is contemplated that the nucleotide sequences presented herein may contain some small percentage of errors. These errors may arise in the normal course of determination of nucleotide sequences. Sequence errors can be corrected by obtaining seeds deposited under the accession numbers cited above, propagating them, isolating genomic DNA or appropriate mRNA from the resulting plants or seeds thereof, amplifying the relevant portion of the genomic DNA or mRNA using primers having a sequence that flanks the erroneous sequence, and sequencing the amplification product.

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#### I.A. Probes, Primers and Substrates

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SDFs of the invention can be applied to substrates for use in array applications such as, but not limited to, assays of global gene expression, for example under varying conditions of development, growth conditions. The arrays can also be used in diagnostic or forensic methods.

Probes and primers of the instant invention will hybridize to a polynucleotide comprising a sequence in SEQ TABLES 1 AND 2. Though many different nucleotide sequences can encode an amino acid sequence, in some instances, the sequences of SEQ TABLES 1 AND 2 are preferred for encoding polypeptides of the invention. However, the sequence of the probes and/or primers of the instant invention need not be identical to those in SEQ TABLES 1 AND 2 or the complements thereof. For example, some variation in probe or primer sequence and/or length can allow additional family members to detected, as well as orthologous genes taxonomically distant related sequences. Similarly probes and/or primers of the invention can include additional nucleotides that serve as a label for detecting the formed duplex or for subsequent cloning purposes.

Probe length will vary depending on the application. For use as PCR primers, probes should be 12-40 nucleotides,

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preferably 18-30 nucleotides long. For use in mapping, probes should be 50 to 500 nucleotides, preferably 100-250 nucleotides long. For Southern hybridizations, probes as long as several kilobases can be used as explained below.

The probes and/or primers can be produced by synthetic procedures such as the triester method of Matteucci et al. *J. Am. Chem. Soc.* 103:3185(1981); or according to Urdea et al. *Proc. Natl. Acad.* 80:7461 (1981) or using commercially available automated oligonucleotide synthesizers.

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# I.B. <u>Methods of Detection and Isolation</u>

### B.1. Hybridization

Probes and/or primers can be used for detection and/or isolation of polynucleotide sequences. Such polynucleotides are included in the subject matter of the invention. Depending on the stringency of the conditions under which these probes and/or primers are used, polynucleotides exhibiting a wide range of similarity to those in SEQ TABLES 1 AND 2 can be detected or isolated.

"Stringency" is a function of probe length, probe composition (G + C content), and salt concentration, organic solvent concentration, and temperature of hybridization or wash conditions. Stringency is typically compared by the parameter is the temperature at which 50% which complementary molecules in the hybridization are hybridized, in terms of a temperature differential from Tm. High stringency conditions are those providing a condition of  $T_m$  -  $5^{\circ}C$  to  $T_m$  -Medium stringency conditions are those providing  $T_{m}$  - $20^{\circ}\text{C}$  to  $T_{m}$  -  $29^{\circ}\text{C}$ . Low stringency conditions are those providing a condition of  $T_m - 40^{\circ}C$  to  $T_m - 48^{\circ}C$ . relationship of hybridization conditions to  $T_m$  (in  ${}^{\circ}C$ ) expressed in the mathematical equation

$$T_{m} = 81.5 - 16.6(\log_{10}[Na^{+}]) + 0.41(%G+C) - (600/N)$$
 (1)

where N is the length of the probe. This equation works well for probes 14 to 70 nucleotides in length that are identical to the target sequence. The equation below for  $T_m$  of DNA-DNA hybrids is useful for probes in the range of 50 to greater than 500 nucleotides, and for conditions that include an organic solvent (formamide).

 $T_m = 81.5 + 16.6 \log \{ [Na^+]/(1 + 0.7 \{Na^+]) \} + 0.41 ( G+C) - 500/L 0.63 ( formamide)$  (2)

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where L is the length of the probe in the hybrid. (P. Tijessen, "Hybridization with Nucleic Acid Probes" in Laboratory Techniques in Biochemistry and Molecular Biology, P.C. vand der Vliet, ed., c. 1993 by Elsevier, Amsterdam.) The T<sub>m</sub> of equation (2) is affected by the nature of the hybrid; for DNA-RNA hybrids T<sub>m</sub> is 10-15°C higher than calculated, for RNA-RNA hybrids T<sub>m</sub> is 20-25°C higher. Because the T<sub>m</sub> decreases about 1 °C for each 1% decrease in homology when a long probe is used (Bonner et al., *J. Mol. Biol.* 81:123 (1973)), stringency conditions can be adjusted to favor detection of identical genes or related family members.

Equation (2) is derived assuming equilibrium and therefore, hybridizations according to the present invention are most preferably performed under conditions of probe excess and for sufficient time to achieve equilibrium. The time required to reach equilibrium can be shortened by inclusion of a "hybridization accelerator" such as dextran sulfate or another high volume polymer in the hybridization buffer.

When the practitioner wishes to examine the result of membrane hybridizations under a variety of stringencies, an efficient way to do so is to perform the hybridization under a low stringency condition, then to wash the hybridization membrane under increasingly stringent conditions. With

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respect to wash steps, preferred stringencies lie within the ranges stated above; high stringency is  $5-8^{\circ}\text{C}$  below  $T_m$ , medium stringency is  $26-29^{\circ}\text{C}$  below  $T_m$  and low stringency is  $45-48^{\circ}\text{C}$  below  $T_m$ .

A number of methods known to those skilled in the art can be used with the probes and/or primers of the invention to isolate and detect polynucleotides, including, without limitation: Southerns, Northerns, Branched DNA hybridization assays, polymerase chain reaction, and variations thereof.

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When using SDFs to identify orthologous genes in other species, the practitioner will preferably adjust the amount of target DNA of each species so that, as nearly as is practical, the same number of genome equivalents are present for each species examined. This prevents faint signals from species having large genomes, and thus small numbers of genome equivalents per mass of DNA, from erroneously being interpreted as absence of the corresponding gene in the genome.

A good general discussion of the factors for determining hybridization conditions is provided by Sambrook et al. ("Molecular Cloning, a Laboratory Manual, 2nd ed., c. 1989 by Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY; see esp., chapters 11 and 12). Additional considerations and details of the physical chemistry of hybridization are provided by G.H. Keller and M.M. Manak "DNA Probes", 2<sup>nd</sup> Ed. pp. 1-25, c. 1993 by Stockton Press, New York, NY.

Hybridization of one nucleic acid to another constitutes a physical property that defines the subject SDF of the invention. Also, such hybridization imposes structural limitations on the pair. For example, for a probe molecule, given that the sequence of the probe nucleic acid is known and fixed, equation (2) indicates that the combined variation in GC content of the target DNA and mismatch between the

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probe and the hybridizing DNA is determined for any given hybridization buffer composition and  $T_{\text{m}}.$ 

The probes and/or primers of the instant invention can be used to detect or isolate nucleotides that are "identical" to the probes or primers. Two nucleic acid sequences or polypeptides are said to be "identical" if the sequence of nucleotides or amino acid residues, respectively, in the two sequences is the same when aligned for maximum correspondence as described below. The term "complementary to" is used herein to mean that the sequence can form a Watson-Crick base pair with a reference polynucleotide sequence. Complementary sequences can include nucleotides, such as inosine, that neither disrupt Watson-Crick base pairing nor contribute to the pairing.

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15 Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Waterman APL. Math. 2:482 (1981), by the homology alignment algorithm of Needleman and Wunsch J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson and Lipman Proc. 20 Natl. Acad. Sci. (USA) 85: 2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, BLAST, PASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group (GCG), 575 Science Dr., Madison, WI), by inspection. Given that two sequences have 25 identified for comparison, GAP and BESTFIT are preferably employed to determine their optimal alignment. Typically, the default values of 5.00 for gap weight and 0.30 for gap weight length are used.

The probes and/or primers of the invention can also be used to detect and/or isolate polynucleotides exhibiting at least 80% sequence identity with the sequences of SEQ TABLES 1 AND 2 or fragments thereof.

"Percentage of sequence identity" is determined by comparing two optimally aligned sequences over a comparison

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window, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (e.g., gaps or overhangs) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity. "Percentage of sequence identity" can be determined by the algorithms described above.

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The term "substantial identity" between polynucleotide or polypeptide sequences refers to polynucleotide or polypeptide comprising a sequence that has at least 80% sequence identity, preferably at least 85%, more preferably at least 90% and most preferably at least 95%, even more preferably, at least 96%, 97%, 98% or 99% sequence identity compared to a reference sequence using the programs.

Isolated polynucleotides within the scope of the invention also include allelic variants of the sequences presented in SEQ TABLES 1 AND 2. An "allelic variant" is a sequence that is a variant from that of the represents the same chromosomal locus but Allelic variants can arise by normal genetic variation in a population. Allelic variants can also be produced by genetic engineering methods. An allelic variant can be one that is found in a naturally occurring plant, including a cultivar or ecotype. A silent allele can give rise to phenotypic and expression profiles. An allelic variant may or may not give rise to a phenotypic change, and may or may not be expressed. An expressed allele can result in a detectable change in the phenotype of the trait

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represented by the locus. Allelic variations can occur in any portion of the gene sequence, including regulatory regions as well as structural regions.

With respect to nucleotide sequences, degeneracy of the genetic code provides the possibility to substitute at least one base of the base sequence of a gene with a different base without causing the amino acid sequence of the polypeptide produced from the gene to be changed. Hence, the DNA of the present invention may also have any base sequence that has been changed from a sequence in SEQ TABLES 1 AND 2 by substitution in accordance with degeneracy of genetic code. References describing codon usage include: Carels et al., J. Mol. Evol. 46: 45 (1998) and Fennoy et al., Nucl. Acids Res. 21(23): 5294 (1993).

## A. B.2. Mapping

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The isolated SDF DNA of the invention can be used to create various types of genetic and physical maps of the genome of corn, Arabidopsis or other plants. Some SDFs may be absolutely associated with particular phenotypic traits, allowing construction of gross genetic maps. While not all SDFs will immediately be associated with a phenotype, all SDFs can be used as probes for identifying polymorphisms associated with phenotypes of interest. Briefly, total DNA is isolated from individuals and is subsequently cleaved with one or more restriction enzymes, separated according to mass, transferred to a solid support, hybridized with SDF DNA and the pattern of fragments compared. Polymorphisms associated with a particular SDF are visualized as differences in the size of fragments produced between individual DNA samples after digestion with a particular restriction enzyme and hybridization with the SDF. After identification of polymorphic SDF sequences, linkage studies can be conducted. Recombinants produced are analyzed using the same restriction

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enzyme/hybridization procedure. After identification of many polymorphisms using SDF sequences, linkage studies can be conducted by using the individuals showing polymorphisms as parents in crossing programs. F2 progeny recombinants or recombinant inbreds, for example, are then analyzed using the 5 same restriction enzyme/hybridization procedure. of DNA polymorphisms along the chromosomes can be inferred based on the frequency with which they are inherited together versus independently. The closer two polymorphisms are 10 together in a chromosome the higher the probability that they are inherited together. Integration of the relative positions of all the polymorphisms and associated marker SDFs produces a genetic map of the species, where the distances between markers reflect the recombination frequencies in that 15 chromosome segment.

The use of recombinant inbred lines for such genetic mapping is described for Arabidopsis by Alonso-Blanco et al. (Methods in Molecular Biology, vol.82, "Arabidopsis Protocols", pp. 137-146, J.M. Martinez-Zapater and J.

20 Salinas, eds., c. 1998 by Humana Press, Totowa, NJ) and for corn by Burr ("Mapping Genes with Recombinant Inbreds", pp. 249-254. In Freeling, M. and V. Walbot (Ed.), The Maize Handbook, c. 1994 by Springer-Verlag New York, Inc.: New York, NY, USA; Berlin Germany; Burr et al. Genetics (1998)

25 118: 519; Gardiner, J. et al., (1993) Genetics 134: 917). However, this procedure is not limited to plants and can be used for other organisms (such as yeast) or for individual cells.

The SDFs of the present invention can also be used for simple sequence repeat (SSR) mapping. Rice SSR mapping is described by Morgante et al. (The Plant Journal (1993) 3: 165), Panaud et al. (Genome (1995) 38: 1170); Senior et al. (Crop Science (1996) 36: 1676), Taramino et al. (Genome (1996) 39: 277) and Ahn et al. (Molecular and General

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Genetics (1993) 241: 483-90). SSR mapping can be achieved using various methods. In one instance, polymorphisms are identified when sequence specific probes flanking an SSR contained within an SDF are made and used in polymerase chain 5 reaction (PCR) assays with template DNA from two or more individuals of interest. Here, a change in the number of tandem repeats between the SSR-flanking sequence produces differently sized fragments (U.S. Patent 5,766,847). Alternatively, polymorphisms can be identified by using the 10 PCR fragment produced from the SSR-flanking sequence specific primer reaction as a probe against Southern blots representing different individuals (U.H. Refseth et al., (1997) Electrophoresis 18: 1519).

Genetic and physical maps of crop species have many uses. For example, these maps can be used to devise positional cloning strategies for isolating novel genes from the mapped crop species. In addition, because the genomes of closely related species are largely syntenic (that is, they display the same ordering of genes within the genome), these maps can be used to isolate novel alleles from wild relatives of crop species by positional cloning strategies.

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The various types of maps discussed above can be used with the SDFs of the invention to identify Quantitative Trait Loci (QTLs). Many important crop traits, such as the solids content of tomatoes, are quantitative traits and result from the combined interactions of several genes. These genes reside at different loci in the genome, oftentimes on different chromosomes, and generally exhibit multiple alleles at each locus. The SDFs of the invention can be used to identify QTLs and isolate specific alleles as described by de Vicente and Tanksley (Genetics 134:585 (1993)). In addition to isolating QTL alleles present crop species, the SDFs of the invention can also be used to isolate alleles from the corresponding QTL of wild relatives. Transgenic plants

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having various combinations of QTL alleles can then be created and the effects of the combinations measured. Once an ideal allele combination has been identified, crop improvement can be accomplished either through biotechnological means or by directed conventional breeding

programs (for review see Tanksley and McCouch, Science 277:1063 (1997)).

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In another embodiment the SDFs can be used to help create physical maps of the genome of corn, Arabidopsis and related species. Where SDFs have been ordered on a genetic map, as described above, then SDFs can be used as probes to discover which clones in large libraries of plant DNA fragments in YACs, BACs, etc. contain the same SDF or similar sequences, thereby facilitating the assignment of the large DNA fragments to chromosomal positions. Subsequently, the large BACs, YACs, etc. can be ordered unambiguously by more detailed studies of their sequence composition (e.g. Marra et al. (1997) Genomic Research 7:1072-1084) and by using their end or other sequences to find the identical sequences in other cloned DNA fragments. The overlapping of DNA sequences in this way allows large contigs of plant sequences to be built, that, when sufficiently extended, provide a complete physical map of a chromosome. Sometimes the SDFs themselves will provide the means of joining cloned sequences into a contig.

The patent publication WO95/35505 and U.S. Patents 5,445,943 and 5,410,270 describe scanning multiple alleles of a plurality of loci using hybridization to arrays of oligonucleotides. These techniques are useful for each of the types of mapping discussed above.

Following the procedures described above and using a plurality of the SDFs of the present invention, any individual can be genotyped. These individual genotypes can be used for the identification of particular cultivars,

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varieties, lines, ecotypes and genetically modified plants or can serve as tools for subsequent genetic studies involving multiple phenotypic traits.

#### B. 3 Southern Blot Hybridization

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The sequences from SEQ TABLES 1 AND 2 can be used as probes for various hybridization techniques. These techniques are useful for detecting target polynucleotides in a sample or for determining whether transgenic plants, seeds or host cells harbor a gene or sequence of interest and thus might be expected to exhibit a particular trait or phenotype.

addition, the hybridization of the SDFs of invention to nucleic acids obtained from other organisms can be used to identify orthologous genes from other species and/or additional members of gene families either in the same or different species. In regard to identifying genes in species, a Southern blot of genomic DNA provides description of isolated DNA fragments that comprise orthologous genes or additional members of the gene families. That is, given such data, one of ordinary skill in the art could distinguish the isolated DNA fragments by their size together with the restriction sites at each end and by the property of hybridizing with the SDF probe under the stated conditions.

In addition, the SDFs from the invention can be used to isolate additional members of gene families from the same species and/or orthologous genes from different species. This is accomplished by hybridizing an SDF to a Southern blot containing the appropriate genomic DNA or cDNA. Given the resulting hybridization data, one of ordinary skill in the art could distinguish and isolate the correct DNA fragments by size, restriction sites and stated hybridization conditions from a gel or from a library.

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Identification and isolation of orthologous genes from closely related species and alleles within a species particularly desirable because of their potential for crop improvement. Many important crop traits, such as the solid content of tomatoes, result from the combined interactions of the products of several genes residing at different loci in the genome. Generally, alleles at each of these loci can make quantitative differences to the trait. By identifying and isolating numerous alleles for each locus from within or different species, transgenic plants having various combinations of alleles can be created and the effects of the combinations measured. Once a more favorable ideal allele combination has been identified, crop improvement can be accomplished either through biotechnological means or by directed conventional breeding programs (Tanksley et Science 277:1063(1997)).

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The results from hybridizations of the SDFs of the invention to Southern blots containing DNA from another species can also be used to generate restriction fragment maps for the corresponding genomic regions. These maps provide map provides additional information about the relative positions of restriction sites within fragments, further distinguishing mapped DNA from the remainder of the genome.

Physical maps can be made by digesting genomic DNA with different combinations of restriction enzymes.

Probes for Southern blotting to distinguish individual restriction fragments can range in size from 15 to 20 nucleotides to several thousand nucleotides. More preferably, the probe is 100 to 1000 nucleotides long for identifying members of a gene family when it is found that repetitive sequences would complicate the hybridization. For identifying an entire corresponding gene in another species, the probe is more preferably the length of the gene,

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typically 2000 to 10,000 nucleotides, but probes 50-1,000 nucleotides long might be used. Some genes, however, might require probes up to 15,000 nucleotides long or overlapping probes constituting the full-length sequence to span their lengths.

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is preferred that the Also. while it probe be homogeneous with respect to its sequence, that is example, as described below, necessary. For representing members of а gene family having diverse sequences can be generated using PCR to amplify genomic DNA or RNA templates using primers derived from SDFs that include sequences that define the gene family.

For identifying corresponding genes in another species, the probe for Southern blotting most preferably would be the genomic copy of the probe gene. This allows all elements of the gene to be identified in the other species. most preferable probe is a cDNA spanning the entire coding sequence, which allows all of the mRNA-coding portion of the gene to be identified; in this case it is possible that some introns in the gene might be missed. Probes for Southern blotting can easily be generated from SDFs by making primers having the sequence at the ends of the SDF and using corn or Arabidopsis genomic DNA as a template. In instances where the SDF includes sequence conserved among species, primers including the conserved sequence can be used for PCR with genomic DNA from a species of interest to obtain a probe. Similarly, if the SDF includes a domain of interest, that portion of the SDF can be used to make primers and, with appropriate template DNA, used to make a probe to identify genes containing the domain. Alternatively, the PCR products can be resolved, for example by gel electrophoresis, and cloned and/or sequenced. In this manner, the variants of the domain among members of a gene family, both within and across species, can be examined.

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#### B.4.1 Isolating DNA from Related Organisms

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The SDFs of the invention can be used to isolate the corresponding DNA from other organisms. Either cDNA or genomic DNA can be isolated. For isolating genomic DNA, a lambda, cosmid, BAC or YAC, or other large insert genomic library from the plant of interest can be constructed using standard molecular biology techniques as described in detail by Sambrook et al. 1989 (Molecular Cloning: A Laboratory Manual, 2<sup>nd</sup> ed. Cold Spring Harbor Laboratory Press, New York) and by Ausubel et al. 1992 (Current Protocols in Molecular Biology, Greene Publishing, New York).

To screen a phage library, recombinant lambda clones are on appropriate bacterial medium using appropriate E. coli host strain. The resulting plaques are lifted from the plates using nylon or nitrocellulose filters. plaque lifts are processed through denaturation, neutralization, and washing treatments following the standard protocols outlined by Ausubel et al. (1992). The plaque lifts are hybridized to either radioactively labeled or nonradioactively labeled SDF DNA at room temperature for about 16 hours, usually in the presence of 50% formamide and 5X SSC (sodium chloride and sodium citrate) buffer and blocking reagents. The plague lifts are then washed at 42°C with 1% Dodecyl Sulfate (SDS) and Sodium at a particular concentration of SSC. The SSC concentration used dependent upon the stringency at which hybridization occurred in the initial Southern blot analysis performed. example, if a fragment hybridized under medium stringency (e.g., Tm - 20°C), then this condition is maintained or preferably adjusted to a less stringent condition (e.g., Tm-30°C) to wash the plague lifts. Positive clones show detectable hybridization e.g., by exposure to X-ray films or chromogen formation. The positive clones are then

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subsequently isolated for purification using the same general protocol outlined above. Once the clone is purified, restriction analysis can be conducted to narrow the region corresponding to the gene of interest. The restriction analysis and succeeding subcloning steps can be done using procedures described by, for example Sambrook et al. (1989) cited above.

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To screen a YAC library, the procedures outlined for the lambda library are essentially similar except the YAC clones are harbored in bacterial colonies. The YAC clones are plated out at reasonable density on nitrocellulose or nylon filters supported by appropriate bacterial medium in petri Following the growth of the bacterial clones, the plates. filters processed through the denaturation, are neutralization, and washing steps following the procedures of Ausubel et al. 1992. The same hybridization procedures for lambda library screening are followed.

To isolate cDNA, similar procedures using appropriately modified vectors are employed. For instance, the library can be constructed in a lambda vector appropriate for cloning cDNA such as  $\lambda$ gtll. Alternatively, the cDNA library can be made in a plasmid vector. cDNA for cloning can be prepared by any of the methods known in the art, but is preferably prepared as described above. Preferably, a cDNA library will include a high proportion of full-length clones.

#### B. 5. Isolating and/or Identifying Orthologous Genes

Probes and primers of the invention can be used to identify and/or isolate polynucleotides related to those in SEQ TABLES 1 AND 2. Related polynucleotides are those that are native to other plant organisms and exhibit either similar sequence or encode polypeptides with similar biological activity. One specific example is an orthologous gene, a gene that has a high degree of sequence similarity, often along the

28 entire length of the coding portion of the gene, and also encodes a gene product that performs a similar function in the Orthologous genes may be distinguished from homologous genes in that homologous genes share sequence similarity but often only in a portion of the sequence, which often represents a functional domain such as a tyrosine kinase activity, a DNA binding domain, or the like. The functional activities of homologous genes are not necessarily the same, but are the same for orthologous genes. The degree of identity is a function of evolutionary separation and, in closely related species, the degree of identity can be 98 to 100%. The amino acid sequence of a protein encoded by an orthologous gene can be less than 75% identical, but tends to be at least75% or at least 80% identical, more preferably at least 90%, most preferably at least 95% identical to the amino acid sequence of the reference protein.

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To find orthologous genes, the probes are hybridized to nucleic acids from a species of interest under low stringency conditions and blots are then washed under conditions of It is preferable that increasing stringency. stringency be such that sequences that are 85 to 100% identical will hybridize. More preferably, sequences 90 to identical will hybridize and most preferably only sequences greater than 95% identical will hybridize. The low stringency condition is preferably one where sequences containing as much as 40-45% mismatches will be able to hybridize. This condition is established by  $T_m$  -  $40^{\circ}$ C to Tm -  $48^{\circ}$ C (see below). ordinary skill in the art will recognize that, due degeneracy in the genetic code, amino acid sequences that are identical can be encoded by DNA sequences as little as 67% Thus, it is preferable to make an overlapping series of shorter probes, on the order of 24 to 45 nucleotides, and individually hybridize them to the same arrayed library to

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avoid the problem of degeneracy introducing large numbers of mismatches.

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As evolutionary divergence increases, genome sequences also tend to diverge. Thus, one of skill will recognize that searches for orthologous genes between more divergent species will require the use of lower stringency conditions compared between closely related searches species. degeneracy is more of a problem for searches in the genome of a species more distant evolutionarily from the species that is the source of the SDF probe sequences.

Therefore the method described in Bouckaert et al., U.S. Ser. No. 60/121,700 Atty. Dkt. No. 2750-117P, Client Dkt. No. 00010.001, filed February 25, 1999, hereby incorporated in its entirety by reference, can be applied to the SDFs of the present invention to isolate related genes from plant species which do not hybridize to the corn or Arabidopsis sequences of SEO TABLES 1 AND 2.

Identification of the relationship of nucleotide or amino acid sequences among plant species can be done by comparison of the subject nucleotide or amino acid sequence to the sequences of SDFs of the present application presented in SEQ TABLES 1 and 2.

The SDFs of the invention can also be used as probes to search for genes that are related to the SDF within a species. Such related genes are typically considered to be members of a "gene family." In such a case, the sequence similarity will often be concentrated into one or a few portions of the sequence. The portions of similar sequence that define the gene family typically encode a portion of a protein or RNA that has an enzymatic or structural function. The degree of identity in the amino acid sequence of the domain that defines the gene family is preferably at least 70%, more preferably 80 to 95%, most preferably 85 to 99%. To search for members of a gene family within a species, a

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"low stringency" hybridization is usually performed, but this will depend upon the size, distribution and degree of sequence divergence of domains that define the gene family. SDFs encompassing regulatory regions can be used to identify "coordinately expressed" genes by using the regulatory region portion of the SDF as a probe.

In the instances where the SDFs are identified as being expressed from genes that confer a particular phenotype, then the SDFs can also be used as probes to assay plants of different species for those phenotypes.

#### I.C. Methods to Inhibit Gene Expression

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In some instances it is desirable to suppress expression of an endogenous or exogenous gene. A well-known instance is the FLAVOR-SAVOR™ tomato, in which the gene encoding ACC synthase is inactivated by an antisense approach, thus delaying softening of the fruit after ripening. See for example, U.S. Patent No. 5,859,330; U.S. No. 5,723,766; Oeller, et al, Science, 254:437-439(1991); Hamilton et al, *Nature*, 346:284-287 (1990). Also, timing of flowering can be controlled by suppression of the FLOWERING LOCUS C; high levels of this transcript are associated with late flowering, while absence of FLC is associated with early flowering (S.D. Michaels et al., Plant Cell 11:949 (1999). Also, the transition of apical meristem from production of leaves with associated shoots to flowering is regulated by TERMINAL FLOWER1, APETALA1 and LEAFY. Thus, when it is desired to induce a transition from shoot production to flowering, it is desirable to suppress *TFL1* expression Liljegren, Plant Cell 11:1007 (1999)). As another instance, it has been found that suppression of the ethylene forming enzyme results in arrested ovule development and female sterility that can be reversed by application of ethylene (D. De Martinis et al., Plant Cell 11:1061 (1999)). The ability

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to manipulate fertility of female plants is useful in the increasing fruit production and creating hybrids.

In an instance when it is desired to express a dominant negative mutation, it often helpful to suppress expression of the endogenous, native protein. Expression of dominant negative mutant proteins is a useful tool for research, for example when a dominant negative mutation of a receptor is used to constitutively activate or suppress a signal transduction cascade, allowing examination of the phenotype and thus the trait(s) controlled by that receptor and pathway.

#### C.1 Antisense

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In the case of polynucleotides used to inhibit expression of an endogenous gene, the introduced sequence need not be perfectly identical to a sequence of the target endogenous gene. The introduced polynucleotide sequence will typically be at least substantially identical (as determined above) to the target endogenous sequence.

Some polynucleotide SDFs in SEQ TABLES 1 AND 2 represent sequences that are expressed in corn and/or Arabidopsis. Thus the invention includes using these sequences to generate antisense constructs to inhibit transcription and/or translation of said SDFs, typically in a plant cell.

To accomplish this, a polynucleotide segment from the desired gene that can hybridize to the mRNA expressed from the desired gene (the "antisense segment") is operably linked to a promoter such that the antisense strand of RNA will be transcribed when the construct is present in a host cell. A regulated promoter can be used in the construct to control transcription of the antisense segment so that transcription occurs only under desired circumstances.

The antisense segment to be introduced generally will be substantially identical to at least a portion of the endogenous gene or genes to be repressed. The sequence, however, need not

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be perfectly identical to inhibit expression. Further, the antisense product may hybridize to the untranslated region instead of or in addition to the coding portion of the gene. The vectors of the present invention can be designed such that the inhibitory effect applies to other proteins within a family of genes exhibiting homology or substantial homology to the target gene.

For antisense suppression, the introduced antisense segment sequence also need not be full length relative to either the primary transcription product or fully processed mRNA. Generally, higher sequence identity can be used to compensate for the use of a shorter sequence. Furthermore, the introduced sequence need not have the same intron or exon pattern, and homology of non-coding segments may be equally effective. Normally, a sequence of between about 30 or 40 nucleotides and the full length of the transcript should be used, though a sequence of at least about 100 nucleotides is preferred, a sequence of at least about 200 nucleotides is preferred, and a sequence of at least 500 about nucleotides is especially preferred.

#### C.2. Ribozymes

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Ιt contemplated that is also gene constructs representing ribozymes and based on the SDFs in SEQ TABLES 1 AND 2 are an object of the invention. Ribozymes can also be used to inhibit expression of genes by suppressing the translation of the mRNA into a polypeptide. It is possible to design ribozymes that specifically pair with virtually any target RNA and cleave the phosphodiester backbone at a specific location, thereby functionally inactivating the target RNA. In carrying out this cleavage, the ribozyme is not itself altered, and is thus capable of recycling and cleaving other molecules, making it a true enzyme. The inclusion of ribozyme sequences

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within antisense RNAs confers RNA-cleaving activity upon them, thereby increasing the activity of the constructs.

A number of classes of ribozymes have been identified. One class of ribozymes is derived from a number of small circular RNAs. which are capable of self-cleavage replication in plants. The RNAs replicate either alone RNAs) or with a helper virus (satellite RNAs). (viroid Examples include RNAs from avocado sunblotch viroid and the satellite RNAs from tobacco ringspot virus, luceme transient streak virus, velvet tobacco mottle virus, solanum nodiflorum mottle virus and subterranean clover mottle virus. design and use of target RNA-specific ribozymes is described in Haselhoff et al. Nature, 334:585 (1988).

the antisense constructs above, the sequence portion necessary for pairing need not be identical to the target nucleotides to be cleaved, nor identical to the sequences in SEQ TABLES 1 AND 2. Generally, the sequence in ribozyme capable of binding to the target sequence exhibits substantial sequence identity to a sequence in SEQ TABLES 1 AND 2 or the complement thereof, or to a portion of said sequence or complement. Further, the ribozyme sequence also need not be full length relative to either the primary transcription product or fully processed mRNA. The ribozyme can be equally effective in inhibiting mRNA translation by cleaving either in the untranslated or coding regions. Generally, higher sequence identity can be used to compensate use of a shorter sequence. Furthermore, the introduced sequence need not have the same intron or exon pattern, and homology of non-coding segments may be equally effective.

#### C.3. Sense Suppression

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Another method of suppression is by introducing an exogenous copy of the gene to be suppressed. Introduction of

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expression cassettes in which a nucleic acid is configured in the sense orientation with respect to the promoter into the chromosome of a plant or by a self-replicating virus has been shown to be an effective means by which to induce degradation of mRNAs of target genes. For an example of the use of this method to modulate expression of endogenous genes see, Napoli et al., The Plant Cell 2:279 (1990), and U.S. Patents Nos. 5,034,323, 5,231,020, and 5,283,184. Generally, inhibition of expression is desired, some transcription of the introduced sequence is probably necessary. The effect may occur where the introduced sequence contains no coding sequence per se, but comprises only intron or untranslated sequences homologous to sequences present in the primary transcript of the endogenous sequence. The introduction of only regulatory promoter sequences can also cause interference with the activity of endogenous promoters possessing the same sequence. Thus, the described SDFs can also be used to transcription. In all of these procedures, the introduced sequence generally will be substantially identical to the endogenous sequence intended to be inactivated. The minimal identity will typically be greater than about 65%, but a higher identity might exert a more effective repression of expression of the endogenous sequences. Sequence identity of more than about 80% is preferred, though about 95% to absolute identity would be most preferred. As with antisense regulation, the effect would likely apply to any other proteins within a similar family of genes exhibiting homology or substantial homology to the suppressing sequence.

#### C.4. Other Methods to Inhibit Gene Expression

Yet another means of suppressing gene expression is to insert a polynucleotide into the gene of interest to disrupt transcription or translation of the gene.

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Low frequency homologous recombination can be used to target a polynucleotide insert to a gene by flanking the polynucleotide insert with sequences that are substantially similar to the gene to be disrupted. Sequences from SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequence thereto can be used for homologous recombination.

In addition, random insertion of polynucleotides into a host cell genome can also be used to disrupt the gene of interest. Azpiroz-Leehan et al., Trends in Genetics 13:152 (1997). In this method, screening for clones from a library containing random insertions is preferred to identifying those that have polynucleotides inserted into the gene of interest. Such screening can be performed using probes and/or primers described above based on sequences from SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequence thereto. The screening can also be performed by selecting clones or R<sub>1</sub> plants having a desired phenotype.

## I.D. Methods of Functional Analysis

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The constructs described in the methods under I.C. above can be used to determine the function of the polypeptide encoded by the gene that is targeted by the constructs.

Down-regulating the transcription and translation of the targeted gene, the host cell or organisms, such as a plant, may produce phenotypic changes as compared to a wild-type cell or organism. In addition, in vitro assays can be used to determine if any biological activity, such as calcium flux, DNA transcription, nucleotide incorporation, etc., are being modulated by the down-regulation of the targeted gene.

Coordinated regulation of sets of genes, e.g., those contributing to a desired polygenic trait, is sometimes necessary to obtain a desired phenotype. SDFs of the invention representing transcription activation and DNA binding domains can be assembled into hybrid transcriptional

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activators. These hybrid transcriptional activators can be used with their corresponding DNA elements (i.e., those bound by the DNA-binding SDFs) to effect coordinated expression of desired genes (J.J. Schwarz et al., Mol. Cell. Biol. 12:266 (1992), A. Martinez et al., Mol. Gen. Genet. 261:546 (1999)).

The SDFs of the invention can also be used in the two-hybrid genetic systems to identify networks of protein-protein interactions (L. McAlister-Henn et al., Methods 19:330 (1999), J.C. Hu et al., Methods 20:80 (2000), M. Golovkin et al., J. Biol. Chem. 274:36428 (1999), K. Ichimura et al., Biochem. Biophys. Res. Comm. 253:532 (1998)). The SDFs of the invention can also be used in various expression display methods to identify important protein-DNA interactions (e.g. B. Luo et al., J. Mol. Biol. 266:479 (1997)).

## I.E. Promoters

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The SDFs of the invention are also useful as structural or regulatory sequences in a construct for modulating the expression of the corresponding gene in a plant or other organism, e.g. a symbiotic bacterium. For example, promoter sequences represented in SEQ TABLES 1 AND 2 can be useful in directing expression of coding sequences either as constitutive promoters or to direct expression in particular cell types, tissues, or organs or in response to environmental stimuli.

The term "promoter" refers to a region ofer sequence determinants located upstream or downstream from the start of transcription and which are involved in recognition and binding of RNA polymerase and other proteins to initiate transcription. A "plant promoter" is a promoter capable of initiating transcription in plant cells and can be used to drive expression of a translated portion of an SDF. Such promoters need not be of plant origin. For example, promoters derived from plant viruses, such as the CaMV35S promoter or from

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Agrobacterium tumefaciens such as the T-DNA promoters, can be plant promoters. A typical example of a constitutive promoter of plant origin is the promoter of the cowpea trypsin inhibitor Typical examples of temporal and/or tissue specific of plant origin that can be used with promoters polynucleotides of the present invention, are: PTA29, which is capable of driving gene expression specifically in tapetum and only during anther development (Koltonow et al., Plant Cell 2:1201 (1990); RCc2 and RCc3, promoters that direct root-specific gene expression in rice (Xu et al., Plant Mol. Biol. 27:237 (1995); TobRB27, a rootspecific promoter from tobacco (Yamamoto et al., Plant Cell 3:371 (1991)).

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By "specific promoters" is meant promoters that have a high preference of driving gene expression in the specified tissue and/or at the specified time during the concerned tissue or organ development. By "high preference" is meant at least 3-fold, preferably 5-fold, more preferably at least 10-fold still more preferably at least 20-fold, 50-fold or 100-fold increase in expression in the desired tissue over the expression in any undesired tissue.

A typical example of an inducible promoter, which can be utilized with the polynucleotides of the present invention, is PARSK1, the promoter from the *Arabidopsis* gene encoding a serine-threonine kinase enzyme, and which promoter is induced by dehydration, abscissic acid and sodium chloride (Wang and Goodman, *Plant J.* 8:37 (1995)).

With respect to the SDFs of the present invention a promoter is likely to be a relatively small portion of sequence located in the first genomic DNA (qDNA) nucleotides upstream from an initial exon identified in a gDNA sequence or initial "ATG" or methionine codon corresponding cDNA or mRNA sequence. Such promoters are more likely to be found in the first 1000 nucleotides upstream of an WO 00/40695

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of initial ATG or methionine codon а CDNA corresponding to a gDNA sequence. In particular, the promoter is usually located upstream of the transcription start site. Such a start site is located at the first exon predicted in the OCKHAM-cDNA predictions. In such an instance, the transcription start site is the first nucleotide of the 5' most exon, if the predictions are in the plus (+) strand, or the 3' most if the in the minus (-) strand. predictions are Alternative transcription start sites may be located between the first nucleotide of the 5' most exon (or the 3' most exon in the minus (-) strand) and the initial ATG or methionine codon in the cDNA sequence. The portions of a particular gDNA sequence that function as a promoter in a plant cell will preferably be found to hybridize at medium or high stringency to gDNA sequences presented in SEQ TABLES 1 AND 2.

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Promoters are generally modular in nature. Short DNA sequences representing binding sites for proteins can be separated from each other by intervening sequences of varying For example, within a particular functional module protein binding sites may be constituted by regions of 5 to 60, preferably 10 to 30, more preferably 10 to 20 nucleotides. such binding sites, there are typically 2 nucleotides that specifically contact amino acids of the nucleic acid binding protein. The protein binding sites are usually separated from each other by 10 to several hundred nucleotides, typically by 15 to 150 nucleotides, often by 20 to 50 nucleotides. DNA binding sites in promoter elements often display dyad symmetry in their sequence. Often elements binding several different proteins, and/or a plurality of sites that bind the same protein, will be combined in a region of 100 to 1000 basepairs.

Elements that have transcription regulatory function can be isolated from their corresponding endogenous gene, or the desired sequence can be synthesized, and recombined in

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constructs to direct expression of a structural gene in a desired tissue-specific, temporal-specific or other desired manner of inducibility or suppression. When hybridizations are performed to identify or isolate elements of a promoter by hybridization to the long sequences presented in SEQ TABLES 1 AND 2, conditions should be adjusted to account for the above-described nature of promoters. For example short probes, constituting the element sought, should be used under low temperature and/or high salt conditions. When long probes, which might include several promoter elements are used, low to medium stringency conditions are preferred when hybridizing to promoters across species.

can consist of "basal promoter" Promoters a functions as a site for assembly of a transcription complex comprising an RNA polymerase, for example RNA polymerase II. A typical transcription complex will include additional factors such as  $TF_{II}B$ ,  $TF_{II}D$ , and  $TF_{II}E$ . Of these,  $TF_{II}D$  appears to be the only one to bind DNA directly. Basal promoters frequently include a "TATA box" element usually located \_rtween 20 and 35 nucleotides upstream from the site of initiation of. transcription. Basal promoters also sometimes include a "CCAAT box" element (typically a sequence CCAAT) and/or a GGGCG sequence, usually located between 40 and 200 nucleotides, preferably 60 to 120 nucleotides, upstream from the start site of transcription.

The promoter might also contain one or more "enhancers" and/or "suppressors" that function as binding sites for additional transcription factors that have the function of modulating the level of transcription with respect to tissue specificity of transcription, transcriptional responses to particular environmental or nutritional factors, and the like.

If a nucleotide sequence of an SDF, or part of the SDF, functions as a promoter or portion of a promoter, then nucleotide substitutions, insertions or deletions that do not

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substantially affect the binding of relevant DNA binding proteins would be considered equivalent to the exemplified It is envisioned that there are nucleotide sequence. instances where it is desirable to decrease the binding of relevant DNA binding proteins to "silence" or "down-regulate" a promoter, or conversely to increase the binding of relevant DNA binding proteins to "enhance" or "up-regulate" a In such instances, polynucleotides representing promoter. changes to the nucleotide sequence of the DNA-protein contact region by insertion of additional nucleotides, changes to identity of relevant nucleotides, including use of chemically-modified bases, or deletion of one or nucleotides are considered encompassed by the present invention.

Promoter function can be assayed by methods known in the art, preferably by measuring activity of a reporter gene operatively linked to the sequence being tested for promoter function. Examples of reporter genes include those encoding luciferase, green fluorescent protein, G.S., neo, cat and bar.

## 20 I.F. UTRs and Junctions

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Polynucleotides comprising untranslated (UTR) sequences and intron/exon junctions are also within the scope of the invention. UTR sequences include introns and 5' or 3' untranslated regions (5' UTRs or 3' UTRs). Portions of the sequences shown in SEQ TABLES 1 AND 2 can comprise UTRs and introns or intron/exon junctions.

These portions of SDFs, especially UTRs, can have regulatory functions related to, for example, translation rate and mRNA stability. Thus, these portions of SDFs can be isolated for use as elements of gene constructs for expression of polynucleotides encoding desired polypeptides.

Introns of genomic DNA segments might also have regulatory functions. Sometimes promoter elements,

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especially transcription enhancer or suppressor elements, are found within introns. Also, elements related to stability of heteronuclear RNA and efficiency of transport to the cytoplasm for translation can be found in intron elements. Thus, these segments can also find use as elements of expression vectors intended for use to transform plants.

Just as with promoters, introns and UTR sequences and intron/exon junctions can vary from those shown in SEQ TABLES 1 AND 2. Such changes from those sequences preferably will not affect the regulatory activity of the UTRs or intron or intron/exon junction sequences on expression, transcription, or translation. However, in some instances, down-regulation of such activity may be desired to modulate traits or phenotypic or *in vitro* activity.

# 15 I.G. Coding Sequences

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Isolated polynucleotides of the invention can include coding sequences that encode polypeptides comprising an amino acid sequence encoded by a sequences in SEQ TABLES 1 AND 2 or an amino acid sequence presented in SEQ TABLES 1 AND 2.

A nucleotide sequence "encodes" a polypeptide if a cell (or a cell free in vitro system) expressing that nucleotide sequence produces a polypeptide having the recited amino acid sequence when the nucleotide sequence is transcribed and the primary transcript is subsequently processed and translated by a host cell (or a cell free in vitro system) harboring the nucleic acid. Thus, an isolated nucleic acid that "encodes" a particular amino acid sequence can be a genomic sequence comprising exons and introns or a cDNA sequence represents the product of splicing thereof. An isolated nucleic acid "encoding an amino acid sequence" also encompasses heteronuclear RNA, which contains sequences that are spliced out during expression, and mRNA, which lacks those sequences.

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Coding sequences can be constructed using chemical synthesis techniques or by isolating coding sequences or by modifying such synthesized or isolated coding sequences as described above.

In addition to encoding the polypeptide sequences of SEQ TABLES 1 AND 2, which are native to corn or *Arabidopsis*, the isolated polynucleotides can be variant polynucleotides that encode mutants, fragments, and fusions of those native proteins. Such polypeptides are described below in part II.

In variant polynucleotides generally, the number of substitutions, deletions or insertions is preferably less than 20%, more preferably less than 15%; even more preferably less than 10%, 5%, 3% or 1% of the number of nucleotides comprising a particularly exemplified sequence. It is generally expected that non-degenerate nucleotide sequence changes that result in 1 to 10, more preferably 1 to 5 and most preferably 1 to 3 amino acid insertions, deletions or substitutions will not greatly affect the function of an encoded polypeptide. The most preferred embodiments are those wherein 1 to 20, preferably 1 to 10, most preferably 1 to 5 nucleotides are added to, deleted from and/or substituted in the sequences specifically disclosed in SEQ TABLES 1 AND 2.

Insertions or deletions in polynucleotides intended to be used for encoding a polypeptide should preserve the reading frame. This consideration is not so important in instances when the polynucleotide is intended to be used as a hybridization probe.

# II. Polypeptides

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Polypeptides within the scope of the invention include both native proteins as well as mutants, fragments, and fusions thereof. Polypeptides of the invention are those encoded by any of the six reading frames of sequences shown

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in SEQ TABLES 1 AND 2, preferably encoded by the three frames reading in the 5' to 3' direction of the sequences as shown.

Native polypeptides include the proteins encoded by the sequences shown in SEQ TABLES 1 AND 2. Such native polypeptides include those encoded by allelic variants.

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Variants, including mutants, will exhibit at least 80% sequence identity to those native polypeptides of SEQ TABLES 1 AND 2. Sequence identity is used for polypeptides as defined above for polynucleotides. More preferably, the variants will exhibit at least 85% sequence identity; even more preferably, at least 90% sequence identity; more preferably at least 95%, 96%, 97%, 98%, or 99% sequence identity. "Fragments" of polypeptide or "portions" of polypeptides will exhibit similar degrees of identity to the relevant portions of the native polypeptide. Fusions will exhibit similar degrees of identity in that portion of the fusion represented by the variant of the native peptide.

Furthermore, variants will exhibit at least one of the functional properties of the native protein. Such properties include, without limitation, protein interaction, DNA interaction, biological activity, immunological activity, receptor binding, signal transduction, transcription activity, growth factor activity, secondary structure, three-dimensional structure, etc. As to properties related to in vitro or in vivo activities, the variants preferably exhibit at least 60% of the activity of the native protein; more preferably at least 70%, even more preferably at least 80%, 85%, 90% or 95% of at least one activity of the native protein.

A type of mutant of the native polypeptides comprises amino acid substitutions. "Conservative substitutions" are preferred to maintain the function or activity of the polypeptide. Such substitutions include conservation of charge, polarity, hydrophobicity, size, etc. For example, one or more amino acid residues within the sequence can be

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substituted with another amino acid of similar polarity that acts as a functional equivalent, for example providing a hydrogen bond in an enzymatic catalysis. Substitutes for an amino acid within an exemplified sequence are preferably made among the members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

Within the scope of sequence identity described above, a polypeptide of the invention may have additional individual amino acids or amino acid sequences inserted into the polypeptide in the middle thereof and/or at the N-terminal and/or C-terminal ends thereof. Likewise, some of the amino acids or amino acid sequences may be deleted from the polypeptide.

## Antibodies

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Isolated polypeptides can be utilized to produce antibodies. Polypeptides of the invention can generally be used, for example, as antigens for raising antibodies by known techniques. The resulting antibodies are useful as reagents for determining the distribution of the antigen protein within the tissues of a plant or within a cell of a plant. The antibodies are also useful for examining the expression level of proteins in various tissues, for example in a wild-type plant or following genetic manipulation of a plant, by methods such as Western blotting.

Antibodies of the present invention, both polyclonal and monoclonal, may be prepared by conventional methods. In

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general, the polypeptides of the invention are first used to immunize a suitable animal, such as a mouse, rat, rabbit, or goat. Rabbits and goats are preferred for the preparation of polyclonal sera due to the volume of serum obtainable, and the availability of labeled anti-rabbit and anti-goat antibodies as detection reagents. Immunization is generally performed by mixing or emulsifying the protein in saline, preferably in an adjuvant such as Freund's complete adjuvant, and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). A dose of  $50-200 \mu g/injection$  is typically sufficient. Immunization is generally boosted 2-6 weeks later with one or more injections of the protein in saline, preferably using Freund's incomplete adjuvant. One alternatively generate antibodies by in vitro immunization using methods known in the art, which for the purposes of this invention is considered equivalent to in vivo immunization.

Polyclonal antisera is obtained by bleeding the immunized animal into a glass or plastic container, incubating the blood at 25°C for one hour, followed by incubating the blood at 4°C for 2-18 hours. The serum is recovered by centrifugation (e.g., 1,000xg for 10 minutes). About 20-50 ml per bleed may be obtained from rabbits.

Monoclonal antibodies are prepared using the method of Kohler and Milstein, Nature 256: 495 (1975), or modification thereof. Typically, a mouse or rat is immunized as described above. However, rather than bleeding the animal to extract serum, the spleen (and optinally several large lymph nodes) is removed and dissociated into single cells. If desired, the spleen cells can be screened (after removal of nonspecifically adherent cells) by applying a cell suspension to a plate, or well, coated with the protein antigen. B-cells expressing membrane-bound immunoglobulin specific for the antigen bind to the plate, and are not rinsed away with the rest of the suspension. Resulting B-cells, or all dissociated spleen

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cells, are then induced to fuse with myeloma cells to form hybridomas, and are cultured in a selective medium (e.g., hypoxanthine, aminopterin, thymidine medium, "HAT"). The resulting hybridomas are plated by limiting dilution, and are assayed for the production of antibodies which bind specifically to the immunizing antigen (and which do not bind to unrelated antigens). The selected Mab-secreting hybridomas are then cultured either *in vitro* (e.g., in tissue culture bottles or hollow fiber reactors), or *in vivo* (as ascites in mice).

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Other methods for sustaining antibody-producing B-cell clones, such as by EBV transformation, are known.

desired, the antibodies (whether polyclonal monoclonal) may be labeled using conventional techniques. Suitable labels include fluorophores, chromophores, radioactive atoms (particularly <sup>32</sup>P and <sup>125</sup>I), electron-dense reagents, enzymes, and ligands having specific binding partners. typically detected by ther activity. For example, horseradish peroxidase is usually detected by its ability to convert 3,3',5,5'-tetramethylbenzidine (TNB) to a blue pigment, quantifiable with a spectrophotometer. "Specific binding partner" refers to a protein capable of binding a ligand molcule with high specificity, as for example in the case of an antigen and a monoclonal antibody specific therefor. specific binding partners include biotin and avidin or streptavidin, IgG and protein A, and the numerous receptorligand couples known in the art. It should be understood that the above description is not meant to categorize the various labels into distinct modes. For example,  $^{125}\text{I}$  may serve as a radioactive label or as an electron-dense reagent. serve as an enzyme or as an antigen for a Mab. Further one may combine various labels for desired effect. For example, Mabs and avidin also require labels in the practice of this invention: thus, one might label a Mab with biotin, and detect

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its presence with avidin labeled with <sup>125</sup>I, or with an anti-biotin Mab labeled with HRP. Other permutations and possibilities will be readily apparent to those of ordinary skill in the art, and are considered as equivalents within the scope of the instant invention.

# In Vitro Applications of Polypeptides

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Some polypeptides of the invention will have enzymatic activities that are useful *in vitro*. For example, the soybean trypsin inhibitor (Kunitz) family is one of the numerous families of proteinase inhibitors. It comprises plant proteins which have inhibitory activity against serine proteinases from the trypsin and subtilisin families, thiol proteinases and aspartic proteinases. Thus, these peptides find *in vitro* use in protein purification protocols and perhaps in therapeutic settings requiring topical application of protease inhibitors.

Delta-aminolevulinic acid dehydratase (EC 4.2.1.24) (ALAD) catalyzes the second step in the biosynthesis of heme, the condensation of two molecules of 5-aminolevulinate to form porphobilinogen. Thus, ALAD proteins can be used as catalysts in synthesis of heme derivatives. Enzymes of biosynthetic pathways generally can be used as catalysts for in vitro synthesis of the compounds representing products of the pathway.

Polypeptides encoded by SDFs of the invention can be engineered to provide purification reagents to identify and purify additional polypeptides that bind to them. This allows one to identify proteins that function as multimers or elucidate signal transduction or metabolic pathways. In the case of DNA binding proteins, the polypeptide can be used in a similar manner to identify the DNA determinants of specific binding (S. Pierrou et al., Anal. Biochem. 229:99 (1995), S.

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Chusacultanachai et al., *J. Biol. Chem.* 274:23591 (1999), Q. Lin et al., *J. Biol. Chem.* 272:27274 (1997)).

# II.A. MUTANTS, FRAGMENTS, AND FUSIONS

Generally, mutants, fragments, or fusions of the polypeptides encoded by the maximum length seuqence(MLS) can exhibit at least one of the activities of the identified domains and/or related polypeptides described in Sections (C) and (D) of REF TABLES 1 and 2 corresponding to the MLS of interest.

# 10 II.A.(1) Mutants

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A type of mutant of the native polypeptides comprises amino acid substitutions. "Conservative substitutions", described above (see II.), are preferred to maintain the function or activity of the polypeptide.

polypeptide. Such substitutions include conservation of charge, polarity, hydrophobicity, size, etc. For example, one or more amino acid residues within the sequence can be substituted with another amino acid of similar polarity that acts as a functional equivalent, for example providing a

acts as a functional equivalent, for example providing a hydrogen bond in an enzymatic catalysis. Substitutes for an amino acid within an exemplified sequence are preferably made among the members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine,

include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

Within the scope of sequence identity described above, a polypeptide of the invention may have additional individual

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amino acids or amino acid sequences inserted into the polypeptide in the middle thereof and/or at the N-terminal and/or C-terminal ends thereof. Likewise, some of the amino acids or amino acid sequences may be deleted from the polypeptide. Amino acid substitutions may also be made in the sequences; conservative substitutions being preferred.

One preferred class of mutants are those that comprise (1) the domain of a MLS encoded polypeptide and/or (2) residues conserved between the MLS encoded polypeptide and related polypeptides of the MLS. For this class of mutants, the MLS encoded polypeptide sequence is changed by insertion, deletion, or substitution at positions flanking the domain and/or conserved residues.

Another class of mutants includes those that comprise a MLS encoded polypeptide sequence that is changed in the domain or conserved residues by a conservative substitution.

Yet another class of mutants includes those that lack one of the *in vitro* activities, or structural features of the MLS encoded polypeptides. One example is dominant negative mutants. Such a mutant may comprise an MLS encoded polypeptide sequence with non-conservative changes in a particular domain or group of conserved residues.

#### II.A.(2) FRAGMENTS

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25 Fragments of particular interest are those that comprise a domain identified for a polypeptide encoded by an MLS of the instant invention and mutants thereof. Also, fragments that comprise at least one region of residues conserved between an MLS encoded polypeptide and its related polypeptides are of great interest. Fragments are sometimes useful as dominant negative mutations.

#### II.A.(3) FUSIONS

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Of interest are chimeras comprising (1) a fragment of the MLS encoded polypeptide or mutants thereof of interest and (2) a fragment of a polypeptide comprising the same domain. For example, an AP2 helix encoded by a MLS of the invention fused to second AP2 helix from ANT protein, which comprises two AP2 helices. The present invention also encompasses fusions of MLS encoded polypeptides, mutants, or fragments thereof fused with related proteins or fragments thereof.

## 10 DEFINITION OF DOMAINS

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The polypeptides of the invention may possess identifying domains as shown in REF TABLES 1 and 2. Domains that can fingerprints or signatures be used to are characterize protein families and/or motifs. fingerprints or signatures can comprise conserved (1) primary sequence, (2) secondary structure, and/or (3) dimensional conformation. Generally, each domain has been associated with either a family of proteins or a motif. Typically, these families and/or motifs have been correlated with specific in-vitro and/or in-vivo activities. A domain can be any length, including the entirety of the sequence of a protein. Detailed descriptions of the domains, associated families and motifs, and correlated activities of polypeptides of the instant invention are described below. Usually, the polypeptides with designated domain(s) exhibit at least one activity that is exhibited by any polypeptide that comprises the same domain(s).

Specific domains within the MLS encoded polypeptides are indicated by the reference REF TABLES 1 and 2. In addition, the domains within the MLS encoded polypeptide can be defined by the region that exhibits at least 70% sequence identity with the consensus sequences listed in the detailed description below of each of the domains.

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The majority of the protein domain descriptions given below are obtained from Prosite,

(http://www.expasy.ch/prosite/), and Pfam,

(http://pfam.wustl.edu/browse.shtml).

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1. (AAA) AAA-protein family signature

A large family of ATPases has been described [1 to 5] whose key feature is that they share a conserved region of about 220 amino acids that contains anATP-binding site. This family is now called AAA, for 'A'TPases 'A'ssociated with diverse cellular 'A'ctivities. The proteins that belong to this family either contain one or two AAA domains. Proteins containing two AAA domains:

- Mammalian and drosophila NSF (N-ethylmaleimide-sensitive fusion protein) and the fungal homolog, SEC18. These proteins are involved in intracellular transport between the endoplasmic reticulum and Golgi, as well as between different Golgi cisternae.
- Mammalian transitional endoplasmic reticulum ATPase

  (previously known as p97 or VCP) which is involved in the transfer of membranes from the endoplasmic reticulum to the golgi apparatus. This protein forms a ring-shaped homooligomer composed of six subunits. The yeast homolog is CDC48 and it may play a role in spindle pole proliferation.
- 20 Yeast protein PAS1, essential for peroxisome assembly and the related protein PAS1 from Pichia pastoris.
  - Yeast protein AFG2.

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- Sulfolobus acidocaldarius protein SAV and Halobacterium salinarium cdcH which may be part of a transduction pathway connecting light to cell division.

Proteins containing a single AAA domain:

- Escherichia coli and other bacteria ftsH (or hflB) protein. FtsH is an ATP-dependent zinc metallopeptidase that seems to degrade the heat-shock sigma-32 factor.
- It is an integral membrane protein with a large cytoplasmic C-terminal domain that contain both the AAA and the protease domains.

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- Yeast protein YME1, a protein important for maintaining the integrity of the mitochondrial compartment. YME1 is also a zinc-dependent protease.

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- Yeast protein AFG3 (or YTA10). This protein also seems to contain a AAA domain followed by a zinc-dependent protease domain.

Subunits from the regulatory complex of the 26S proteasome [6] which is involved in the ATP-dependent degradation of ubiquitinated proteins:

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- a) Mammalian subunit 4 and homologs in other higher eukaryotes, in yeast (gene YTA5) and fission yeast (gene mts2).
  - b) Mammalian subunit 6 (TBP7) and homologs in other higher eukaryotes and in yeast (gene YTA2).
  - c) Mammalian subunit 7 (MSS1) and homologs in other higher eukaryotes and in yeast (gene CIM5 or YTA3).
  - d) Mammalian subunit 8 (P45) and homologs in other higher eukaryotes and in yeast (SUG1 or CIM3 or TBY1) and fission yeast (gene let1).

Other probable subunits such as human TBP1 which seems to influences HIV gene expression by interacting with the virus tat transactivator protein and yeast YTA1 and YTA6.

- Yeast protein BCS1, a mitochondrial protein essential for the expression of the Rieske iron-sulfur protein.
- Yeast protein MSP1, a protein involved in intramitochondrial sorting of proteins.
- Yeast protein PAS8, and the corresponding proteins PAS5 from Pichia pastoris and PAY4 from Yarrowia lipolytica.
- Mouse protein SKD1 and its fission yeast homolog (SpAC2G11.06).
  - Caenorhabditis elegans meiotic spindle formation protein mei-1.
  - Yeast protein SAP1.

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- Yeast protein YTA7.

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- Mycobacterium leprae hypothetical protein A2126A.

It is proposed that, in general, the AAA domains in these proteins act as ATP- dependent protein clamps [5]. In addition to the ATP-binding 'A' and 'B' motifs, which are located in the N-terminal half of this domain, there is a highly conserved region located in the central part of the domain which was used to develop a signature pattern.

- Consensus pattern: [LIVMT]-x-[LIVMT]-[LIVMF]-x-[GATMC]-[ST]-[NS]-x(4)-[LIVM]-D-x-A-[LIFA]-x-R
  - [1] Froehlich K.-U., Fries H.W., Ruediger M., Erdmann R., Botstein D., Mecke D. J. Cell Biol. 114:443-453(1991).
- 15 [2] Erdmann R., Wiebel F.F., Flessau A., Rytka J., Beyer A., Froehlich K.-U., Kunau W.-H. Cell 64:499-510(1991).
  - [3] Peters J.-M., Walsh M.J., Franke W.W. EMBO J. 9:1757-1767(1990).
- [4] Kunau W.-H., Beyer A., Goette K., Marzioch M., Saidowsky

  J., Skaletz-Rorowski A., Wiebel F.F. Biochimie 75:209
  224(1993).
  - [5] Confalonieri F., Duguet M. BioEssays 17:639-650(1995).[
  - 6] Hilt W., Wolf D.H. Trends Biochem. Sci. 21:96-102(1996).
- 25 2. Aminotransferases class-IV signature

Aminotransferases share certain mechanistic features with other pyridoxal-phosphate dependent enzymes, such as the covalent binding of the pyridoxal-phosphate group to a lysine residue. On the basis of sequence similarity, these various enzymes can be grouped [1,2] into subfamilies. One of these, called class-IV, currently consists of the following enzymes:

- Branched-chain amino-acid aminotransferase (EC <a href="2.6.1.42">2.6.1.42</a>) (transaminase B), a bacterial (gene ilvE) and eukaryotic enzyme which catalyzes the reversible

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transfer of an amino group from 4-methyl-2oxopentanoate to glutamate, to form leucine and 2oxoglutarate.

- D-alanine aminotransferase (EC <u>2.6.1.21</u>). A bacterial enzyme which catalyzes the transfer of the amino group from D-alanine (and other D-amino acids) to 2-oxoglutarate, to form pyruvate and D-aspartate.
- 4-amino-4-deoxychorismate (ADC) lyase (gene pabC). A bacterial enzyme that converts ADC into 4-aminobenzoate (PABA) and pyruvate.

The above enzymes are proteins of about 270 to 415 amino-acid residues that share a few regions of sequence similarity. Surprisingly, the best-conserved region does not include the lysine residue to which the pyridoxal-phosphategroup is known to be attached, in ilvE. The region that has been selected as a signature pattern is located some 40 residues at the C-terminus side of the PlP-lysine

Consensus pattern: E-x-[STAGCI]-x(2)-N-[LIVMFAC]-[FY]-x(6,12)-[LIVMF]-x-T-x(6,8)-[LIVM]-x-[GS]-[LIVM]-x-[KR]-x(6,12)-[LIVMF]-x-T-x(6,8)-[LIVM]-x-[GS]-[LIVM]-x-[KR]-x(6,12)-[LIVMF]-x-[KR]-x-

- [1] Green J.M., Merkel W.K., Nichols B.P. J. Bacteriol. 174:5317-5323(1992).
- [2] Bairoch A. Unpublished observations (1992).

3. Bacterial mutT domain signature

The bacterial mutT protein is involved in the GO system [1] responsible for removing an oxidatively damaged form of guanine (8-hydroxyguanine or7,8-dihydro-8-oxoguanine) from

DNA and the nucleotide pool. 8-oxo-dGTP is inserted opposite to dA and dC residues of template DNA with almost equal efficiency thus leading to A.T to G.C transversions. MutT specifically degrades 8-oxo-dGTP to the monophosphate with

the concomitant release of pyrophosphate. MutT is a small

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protein of about 12 to 15 Kd. It has been shown [2,3] that a region of about 40 amino acid residues, which is found in the N-terminal part of mutT, can also be found in a variety of other prokaryotic, viral, and eukaryotic proteins. These proteins are:

- Streptomyces pneumoniae mutX.

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- A mutT homolog from plasmid pSAM2 of Streptomyces ambofaciens.
- Bartonella bacilliformis invasion protein A (gene invA).
- Escherichia coli dATP pyrophosphohydrolase.
- Protein D250 from African swine fever viruses.
- Proteins D9 and D10 from a variety of poxviruses.
- Mammalian 7,8-dihydro-8-oxoguanine triphosphatase (EC 3.1.6.-) [4].
- Mammalian diadenosine 5',5'''-P1,P4-tetraphosphate asymmetrical hydrolase (Ap4Aase) (EC 3.6.1.17) [5], which cleaves A-5'-PPPP-5'A to yield AMP and ATP.
- A protein encoded on the antisense RNA of the basic fibroblast growth factor gene in higher vertebrates.
- Yeast protein YSA1.
- Escherichia coli hypothetical protein yfaO.
- Escherichia coli hypothetical protein ygdU and HI0901, the corresponding Haemophilus influenzae protein.
- Escherichia coli hypothetical protein yjaD and HI0432, the corresponding Haemophilus influenzae protein.
  - Escherichia coli hypothetical protein yrfE.
  - Bacillus subtilis hypothetical protein yqkG.
  - Bacillus subtilis hypothetical protein yzgD.
- Yeast hypothetical protein YGL067w.

It is proposed [2] that the conserved domain could be involved in the active center of a family of pyrophosphate-releasing NTPases. As a signature pattern the core region of

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the domain was selected; it contains four conserved glutamate residues.

Consensus pattern: G-x(5)-E-x(4)-[STAGC]-[LIVMAC]-x-R-E-5 [LIVMFT]-x-E-E-

- [1] Michaels M.L., Miller J.H. J. Bacteriol. 174:6321-6325(1992).
- [2] Koonin E.V. Nucleic Acids Res. 21:4847-4847(1993).
- 10 [3] Mejean V., Salles C., Bullions M.J., Bessman M.J., Claverys J.-P. Mol. Microbiol. 11:323-330(1994).
  - [4] Sakumi K., Furuichi M., Tsuzuki T., Kakuma T., Kawabata S., Maki H., Sekiguchi M. J. Biol. Chem. 268:23524-23530(1993).
- 15 [5] Thorne N.M.H., Hankin S., Wilkinson M.C., Nunez C., Barraclough R., McLennan A.G. Biochem. J. 311:717-721(1995).

## 4. Cystatin domain

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This is a very diverse family. Attempts to define separate subfamilies have failed. Typically, either the N-terminal or C-terminal end is very divergent. But splitting into two domains would make very short families. Cathelicidins are related to this family but have not been included. Number of members: 147

- Inhibitors of cysteine proteases [1,2,3], which are found in the tissues and body fluids of animals, in the larva of the worm Onchocerca volvulus [4], as well as in plants, can be grouped into three distinct but related families:
  - Type 1 cystatins (or stefins), molecules of about 100 amino acid residues with neither disulfide bonds nor carbohydrate groups.
  - Type 2 cystatins, molecules of about 115 amino acid residues which contain one or two disulfide loops near their C-terminus.

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- Kininogens, which are multifunctional plasma glycoproteins.

They are the precursor of the active peptide bradykinin and play a role in blood coagulation by helping to position optimally prekallikrein and factor XI next to factor XII. They are also inhibitors of cysteine proteases. Structurally, kininogens are made of three contiguous type-2 cystatin domains, followed by an additional domain (of variable length) which contains the sequence of bradykinin. The first of the three cystatin domains seems to have lost its inhibitory activity.

In all these inhibitors, there is a conserved region of five residues which has been proposed to be important for the binding to the cysteine proteases. The consensus pattern starts one residue before this conserved region.

-Consensus pattern: [GSTEQKRV]-Q-[LIVT]-[VAF]-[SAGQ]-G-x-[LIVMNK]-x(2)-[LIVMFY]-x-[LIVMFYA]-[DENQKRHSIV]

- 20 [1] Barrett A.J. Trends Biochem. Sci. 12:193-196(1987).
  - [2] Rawlings N.D., Barrett A.J. J. Mol. Evol. 30:60-71(1990).
  - [3] Turk V., Bode W. FEBS Lett. 285:213-219(1991).
  - [4] Lustigman S., Brotman B., Huima T., Prince A.M. Mol. Biochem. Parasitol. 45:65-76(1991).

5. Dehydrins signatures

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A number of proteins are produced by plants that experience water-stress. Water-stress takes place when the water available to a plant falls below a critical level. The plant hormone abscisic acid (ABA) appears to modulate the response of plant to water-stress. Proteins that are expressed during water-stress are called dehydrins [1,2] or LEA group 2 proteins [3]. The proteins that belong to this family are listed below.

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- Arabidopsis thaliana XERO 1, XERO 2 (LTI30), RAB18, ERD10 (LTI45) ERD14 and COR47.
- Barley dehydrins B8, B9, B17, and B18.
- Cotton LEA protein D-11.
- 5 Craterostigma plantagineum dessication-related proteins A and B.
  - Maize dehydrin M3 (RAB-17).
  - Pea dehydrins DHN1, DHN2, and DHN3.
  - Radish LEA protein.
- Rice proteins RAB 16B, 16C, 16D, RAB21, and RAB25.
  - Tomato TAS14.
  - Wheat dehydrin RAB 15 and cold-shock protein cor410, cs66 and cs120.

Dehydrins share a number of structural features.

One of the most notable features is the presence, in their central region, of a continuous run of five to nine serines followed by a cluster of charged residues. Such a region has been found in all known dehydrins so far with the exception of pea dehydrins. A second conserved feature is the presence of two copies of alysine-rich octapeptide; the first copy is located just after the cluster of charged residues that follows the poly-serine region and the second copy is found at the C-terminal extremity. Signature patterns for both regions were derived.

Consensus pattern: S(5)-[DE]-x-[DE]-G-x(1,2)-G-x(0,1)-[KR](4)Consensus pattern: [KR]-[LIM]-K-[DE]-K-[LIM]-P-G-

- 30 [1] Close T.J., Kortt A.A., Chandler P.M. Plant Mol. Biol. 13:95-108(1989).
  - [2] Robertson M., Chandler P.M. Plant Mol. Biol. 19:1031-1044(1992).

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[3] Dure L. III, Crouch M., Harada J., Ho T.-H. D., Mundy J., Quatrano R., Thomas T., Sung Z.R. Plant Mol. Biol. 12:475-486(1989).

5 6. D-isomer specific 2-hydroxyacid dehydrogenases (2 Hacid DH)

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This Pfam covers the Formate dehydrogenase, D-glycerate dehydrogenase and D-lactate dehydrogenase families in SCOP. A number of NAD-dependent 2-hydroxyacid dehydrogenases which seem to be specific for the D-isomer of their substrate have been shown [1,2,3,4] to be functionally and structurally related. These enzymes are listed below.

- D-lactate dehydrogenase (EC 1.1.1.28), a bacterial enzyme which catalyzes the reduction of D-lactate to pyruvate.
- D-glycerate dehydrogenase (EC 1.1.1.29) (NADH-dependent hydroxypyruvate reductase), a plant leaf peroxisomal enzyme that catalyzes the reduction of hydroxypyruvate to glycerate. This reaction is part of the glycolate pathway of photorespiration.
- D-glycerate dehydrogenase from the bacteria Hyphomicrobium methylovorum and Methylobacterium extorquens.
- 3-phosphoglycerate dehydrogenase (EC 1.1.1.95), a bacterial enzyme that catalyzes the oxidation of D-3-phosphoglycerate to 3-phosphohydroxypyruvate. This reaction is the first committed step in the 'phosphorylated' pathway of serine biosynthesis.
- Erythronate-4-phosphate dehydrogenase (EC 1.1.1.-) (gene pdxB), a bacterial enzyme involved in the biosynthesis of pyridoxine (vitamin B6).
- D-2-hydroxyisocaproate dehydrogenase (EC 1.1.1.-) (D-hicDH), a bacterial enzyme that catalyzes the reversible and stereospecific interconversion between

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2-ketocarboxylic acids and D-2-hydroxy-carboxylic acids.

- Formate dehydrogenase (EC 1.2.1.2) (FDH) from the bacteria Pseudomonas sp. 101 and various fungi [5].
- Vancomycin resistance protein vanH from Enterococcus faecium; this protein is a D-specific alpha-keto acid dehydrogenase involved in the formation of a peptidoglycan which does not terminate by D-alanine thus preventing vancomycin binding.
- 10 Escherichia coli hypothetical protein ycdW.

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- Escherichia coli hypothetical protein yiaE.
- Haemophilus influenzae hypothetical protein HI1556.
- Yeast hypothetical protein YER081w.
- Yeast hypothetical protein YIL074w.
- All these enzymes have similar enzymatic activities and are structurally related. Three of the most conserved regions of these proteins have been selected to develop patterns. The first pattern is based on a glycine-rich region located in the central section of these enzymes; this region probably corresponds to the NAD-binding domain. The two other patterns contain a number of conserved charged residues, some of which may play a role in the catalytic mechanism.
  - -Consensus pattern: [LIVMA]-[AG]-[IVT]-[LIVMFY]-[AG]-x-G[NHKRQGSAC]-[LIV]-G-x(13,14)-[LIVfMT]-x(2)-[FYwCTH]-[DNSTK]
    -Consensus pattern: [LIVMFYWA]-[LIVFYWC]-x(2)-[SAC]-[DNQHR][IVFA]-[LIVF]-x-[LIVF]-[HNI]-x-P-x(4)-[STN]-x(2)-[LIVMF]-x[GSDN]
- -Consensus pattern: [LMFATC]-[KPQ]-x-[GSTDN]-x-[LIVMFYWR]-30 [LIVMFYW](2)-N-x-[STAGC]-R-[GP]-x-[LIVH]-[LIVMC]-[DNV]
  - [1] Grant G.A. Biochem. Biophys. Res. Commun. 165:1371-1374(1989).

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- [2] Kochhar S., Hunziker P., Leong-Morgenthaler P.M., Hottinger H. Biochem. Biophys. Res. Commun. 184:60-66(1992).
- [3] Ohta T., Taguchi H. J. Biol. Chem. 266:12588-12594(1991).
- [4] Goldberg J.D., Yoshida T., Brick P. J. Mol. Biol.
- 5 236:1123-1140(1994).
  - [5] Popov V.O., Lamzin V.S. Biochem. J. 301:625-643(1994).
  - 7. dnaJ domains signatures and profile

The prokaryotic heat shock protein dnaJ interacts with

the chaperone hsp70-like dnaK protein [1]. Structurally, the
dnaJ protein consists of an N- terminal conserved domain

(called 'J' domain) of about 70 amino acids, a glycine-rich

region ('G' domain') of about 30 residues, a central domain

containing four repeats of a CXXCXGXG motif ('CRR' domain)

and a C-terminal region of 120 to 170 residues. Such a

structure is shown in the following schematic representation:

+-----+ | N-terminal | | Gly-R | | CXXCXGXG | C-terminal | +-----

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It has been shown [2] that the 'J' domain as well as the 'CRR' domain are also found in other prokaryotic and eukaryotic proteins which are listed below.

- a) Proteins containing both a 'J' and a 'CRR' domain:
- Yeast protein MAS5/YDJ1 which seems to be involved in mitochondrial protein import.
  - Yeast protein MDJ1, involved in mitochondrial biogenesis and protein folding.
  - Yeast protein SCJ1, involved in protein sorting.
  - Yeast protein XDJ1.

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- Plants dnaJ homologs (from leek and cucumber).
- Human HDJ2, a dnaJ homolog of unknown function.
- Yeast hypothetical protein YNL077w.
- b) Proteins containing a 'J' domain without a 'CRR' domain:

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- Rhizobium fredii nolC, a protein involved in cultivarspecific nodulation of soybean.

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- Escherichia coli cbpA [3], a protein that binds curved DNA.
- Yeast protein SEC63/NPL1, important for protein assembly into the endoplasmic reticulum and the nucleus.
- Yeast protein SIS1, required for nuclear migration during mitosis.
- 10 Yeast protein CAJ1.

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- Yeast hypothetical protein YFR041c.
- Yeast hypothetical protein YIR004w.
- Yeast hypothetical protein YJL162c.
- Plasmodium falciparum ring-infected erythrocyte

  surface antigen (RESA). RESA, whose function is not known, is associated with the membrane skeleton of newly invaded erythrocytes.
  - Human HDJ1.
  - Human HSJ1, a neuronal protein.
- 20 Drosophila cysteine-string protein (csp).

A signature pattern for the 'J' domain was developed, based on conserved positions in the C-terminal half of this domain. A pattern for the 'CRR' domain, based on the first two copies of that motif was also developed. A profile for the 'J' domain was also developed.

Consensus pattern: [FY]-x(2)-[LIVMA]-x(3)-[FYWHNT]-[DENQSA]-x-L-x-[DN]-x(3)-[KR]-x(2)-[FYI]-

Consensus pattern: C-[DEGSTHKR]-x-C-x-G-x-[GK]-[AGSDM]-x(2)-[GSNKR]-x(4,6)-C-x(2,3)-C-x-G-x-G-

[1] Cyr D.M., Langer T., Douglas M.G. Trends Biochem. Sci. 19:176-181(1994).

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[2] Bork P., Sander C., Valencia A., Bukau B. Trends Biochem. Sci. 17:129-129(1992).

[3] Ueguchi C., Kaneda M., Yamada H., Mizuno T. Proc. Natl. Acad. Sci. U.S.A. 91:1054-1058(1994).

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- 8. Domain of unknown function
- 9. Gamma-thionins family signature

The following small plant proteins are evolutionary

10 related:

- Gamma-thionins from wheat endosperm (gammapurothionins) and barley (gamma- hordothionins) which are toxic to animal cells and inhibit protein synthesis in cell free systems [1].
- A flower-specific thionin (FST) from tobacco [2].
  - Antifungal proteins (AFP) from the seeds of Brassicaceae species such as radish, mustard, turnip and Arabidopsis thaliana [3].
  - Inhibitors of insect alpha-amylases from sorghum [4].
- Probable protease inhibitor P322 from potato.
  - A germination-related protein from cowpea [5].
  - Anther-specific protein SF18 from sunflower [6]. SF18 is a protein that contains a gamma-thionin domain at its N-terminus and a proline-rich C- terminal domain.
  - Soybean sulfur-rich protein SE60 [7].
    - Vicia faba antibacterial peptides fabatin-1 and -2.

In their mature form, these proteins generally consist of about 45 to 50amino-acid residues. As shown in the following schematic representation, these peptides contain eight conserved cysteines involved in disulfide bonds.

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'C': conserved cysteine involved in a disulfide bond.

'\*': position of the pattern.

Consensus pattern: [KRG]-x-C-x(3)-[SV]-x(2)-[FYWH]-x-[GF]-x-5 C-x(5)-C-x(3)-C [The four C's are involved in disulfide bonds]-

- [1] Bruix M., Jimenez M.A., Santoro J., Gonzalez C., Colilla F.J., Mendez E., Rico M. Biochemistry 32:715-724(1993).
- 10 [2] Gu Q., Kawata E.E., Morse M.-J., Wu H.-M., Cheung A.Y. Mol. Gen. Genet. 234:89-96(1992).
  - [3] Terras F.R.G., Torrekens S., van Leuven F., Osborn R.W., Vanderleyden J., Cammue B.P.A., Broekaert W.F. FEBS Lett. 316:233-240(1993).
- 15 [4] Bloch C. Jr., Richardson M. FEBS Lett. 279:101-104(1991).
  - [5] Ishibashi N., Yamauchi D., Miniamikawa T. Plant Mol. Biol. 15:59-64(1990).
  - [7] Choi Y., Choi Y.D., Lee J.S. Plant Physiol. 101:699-700(1993).

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10. haloacid dehalogenase-like hydrolase

This family is structurally different from the alpha/beta hydrolase family (abhydrolase). This family includes L-2-haloacid dehalogenase, epoxide hydrolases and phosphatases.

- 25 The structure of the family consists of two domains. One is an inserted four helix bundle, which is the least well conserved region of the alignment, between residues 16 and 96 of <a href="Swiss:P24069">Swiss:P24069</a>. The rest of the fold is composed of the core alpha/beta domain.
- 30 [1] Hisano T, Hata Y, Fujii T, Liu JQ, Kurihara T, Esaki N, Soda K, J Biol Chem 1996; 271:20322-20330.
  - 11. Helix-turn-helix (HTH3)

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This large family of DNA binding helix-turn helix proteins includes Cro Swiss: P03036 and CI Swiss: P03034.

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12. Heme-binding domain in cytochrome b5 and oxidoreductases (heme 1)

Cytochrome b5 is a membrane-bound hemo protein which acts as an electron carrier for several membrane-bound oxygenases [1]. There are two homologous forms of b5, one found in microsomes and one found in the outer membrane of mitochondria. Two conserved histidine residues serve as axial ligands for the heme group. The structure of a number of oxidoreductases consists of the juxtaposition of a hemebinding domain homologous to that of b5 and either a flavodehydrogenase or a molybdopterin domain. These enzymes are:

- Lactate dehydrogenase (EC  $\underline{1.1.2.3}$ ) [2], an enzyme that consists of a flavodehydrogenase domain and a hemebinding domain called cytochrome b2.
- Nitrate reductase (EC <u>1.6.6.1</u>), a key enzyme involved in the first step of nitrate assimilation in plants, fungi and bacteria [3,4]. Consists of a molybdopterin domain (see <<u>PDOC00484</u>>), a heme-binding domain called cytochrome b557, as well as a cytochrome reductase domain.
- Sulfite oxidase (EC <u>1.8.3.1</u>) [5], which catalyzes the terminal reaction in the oxidative degradation of sulfur-containing amino acids. Also consists of a molybdopterin domain and a heme-binding domain.

This family of proteins also includes:

- TU-36B, a Drosophila muscle protein of unknown function [6].
  - Fission yeast hypothetical protein SpAC1F12.10c.
  - Yeast hypothetical protein YMR073c.
  - Yeast hypothetical protein YMR272c.

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A segment was used which includes the first of the two histidine heme ligands, as a signature pattern for the hemebinding domain of cytochrome b5 family.

- 5 Consensus pattern: [FY]-[LIVMK]-x(2)-H-P-[GA]-G [H is a heme axial ligand]-
  - [1] Ozols J. Biochim. Biophys. Acta 997:121-130(1989).
  - [2] Guiard B. EMBO J. 4:3265-3272(1985).
- 10 [3] Calza R., Huttner E., Vincentz M., Rouze P., Galangau F., Vaucheret H., Cherel I., Meyer C., Kronenberger J., Caboche M. Mol. Gen. Genet. 209:552-562(1987).
  - [4] Crawford N.M., Smith M., Bellissimo D., Davis R.W. Proc. Natl. Acad. Sci. U.S.A. 85:5006-5010(1988).
- 15 [5] Guiard B., Lederer F. Eur. J. Biochem. 100:441-453(1979).
  - [6] Levin R.J., Boychuk P.L., Croniger C.M., Kazzaz J.A., Rozek C.E. Nucleic Acids Res. 17:6349-6367(1989).

# 13. KH domain

- 20 KH motifs probably bind RNA directly. Auto antibodies to Nova, a KH domain protein, cause paraneoplastic opsoclonus ataxia.
  - [1] Burd CG, Dreyfuss G, Science 1994;265:615-621.
  - [2] Musco G, Stier G, Joseph C, Castiglione Morelli MA,
- 25 Nilges M, Gibson TJ, Pastore A, Cell 1996;85:237-245.
  - 14. MAPEG family (aka: FLAP/GST2/LTC4S family signature)

    The following mammalian proteins are evolutionary related [1]:
- Leukotriene C4 synthase (EC 2.5.1.37) (gene LTC4S), an enzyme that catalyzes the production of LTC4 from LTA4.

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- Microsomal glutathione S-transferase II (EC 2.5.1.18) (GST-II) (gene GST2), an enzyme that can also produces LTC4 from LTA4.
- 5-lipoxygenase activating protein (gene FLAP), a protein that seems to be required for the activation of 5-lipoxygenase.

These are proteins of 150 to 160 residues that contain three transmembrane segments. As a signature pattern, a conserved region between the first and second transmembrane domains was selected.

Consensus patternc: G-x(3)-F-E-R-V-[FY]-x-A-[NQ]-x-N-C

- [1] Jakobsson P.-J., Mancini J.A., Ford-Hutchinson A.W. J. Biol. Chem. 271:22203-22210(1996).
- 15. Pathogenesis-related protein Bet v I family signature
  A number of plant proteins, which all seem to be
  involved in pathogen defense response, a.e structurally
  related [1,2,3]. These proteins are:
  - Bet v I, the major pollen allergen from white birch. Bet v I is the main cause of type I allergic reactions in Europe, North America and USSR.
  - Aln g I, the major pollen allergen from alder.
- 25 Api G I, the major allergen from celery.
  - Car b I, the major pollen allergen from hornbeam.
  - Cor a I, the major pollen allergen from hazel.
  - Mal d I, the major pollen allergen from apple.
  - Asparagus wound-induced protein AoPR1.
- Kidney bean pathogenesis-related proteins 1 and 2.
  - Parsley pathogenesis-related proteins PR1-1 and PR1-3.
  - Pea disease resistance response proteins pI49, pI176 and DRRG49-C.
  - Pea abscisic acid-responsive proteins ABR17 and ABR18.

- Potato pathogenesis-related proteins STH-2 and STH-21.
- Soybean stress-induced protein SAM22.

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These proteins are thought to be intracellularly located. They contain from 155 to 160 amino acid residues. As a signature pattern, a conserved region located in the third quarter of these proteins has been selected Consensus pattern: G-x(2)-[LIVMF]-x(4)-E-x(2)-[CSTAEN]-x(8,9)-[GND]-G-[GS]-[CS]-x(2)-K-x(4)-[FY]-

- [1] Breiteneder H., Pettenburger K., Bito A., Valenta R., Kraft D., Rumpold H., Scheiner O., Breitenbach M. EMBO J. 8:1935-1938(1989).
  - [2] Crowell D., John M.E., Russell D., Amasino R.M. Plant Mol. Biol. 18:459-466(1992).
- 15 [3] Warner S.A.J., Scott R., Draper J. Plant Mol. Biol. 19:555-561(1992).
- 16. Photosystem I psaG / psaK (PSI PSAK) proteins signature Photosystem I (PSI) [1] is an integral membrane protein complex that uses light energy to mediate electron transfer from plastocyanin to ferredoxin. It is found in the chloroplasts of plants and cyanobacteria. PSI is composed of at least 14 different subunits, two of which PSI-G (gene psaG) and PSI-K (gene psaK) are small hydrophobic proteins of about 7 to 9 Kd and evolutionary related [2]. Both seem to contain two transmembrane regions. Cyanobacteria seem to encode only for PSI-K.

As a signature pattern, the best-conserved region was selected which seems to correspond to the second transmembrane region.

-Consensus pattern: [GT]-F-x-[LIVM]-x-[DEA]-x(2)-[GA]-x-[GTA]-[SA]-x-[GTA]-[GA]

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- [1] Golbeck J.H. Biochim. Biophys. Acta 895:167-204(1987).
- [2] Kjaerulff S., Andersen B., Nielsen V.S., Moller B.L., Okkels J.S. J. Biol. Chem. 268:18912-18916(1993).
- 17. Plant lipid transfer protein family signature (LTP)

  Plant cells contain proteins, called lipid transfer

  proteins (LTP) [1,2,3], which are able to facilitate the

  transfer of phospholipids and other lipidsacross membranes.

  These proteins, whose subcellular location is not yet known,

  could play a major role in membrane biogenesis by conveying

  phospholipids such as waxes or cutin from their site of

  biosynthesis to membranes unable to form these lipids. Plant

  LTP's are proteins of about 9 Kd (90 amino acids) which

  contain eight conserved cysteine residues all involved in

  disulfide bridges, as shown in the following schematic

\*CxxxxCxxxxxCCxxxxxxxxxxCxCxxxxxxxCxx | | | +----

'C': conserved cysteine involved in a disulfide bond.

'\*': position of the pattern.

representation.

Consensus pattern: [LIVM]-[PA]-x(2)-C-x-[LIVM]-x-[LIVM]-x-25 [LIVMFY]-x-[LIVM]-[ST]-x(3)-[DN]-C-x(2)-[LIVM] [The two C's are involved in disulfide bonds]

- [1] Wirtz K.W.A. Annu. Rev. Biochem. 60:73-99(1991).
- [2] Arondel V., Kader J.C. Experientia 46:579-585(1990).
- 30 [3] Ohlrogge J.B., Browse J., Somerville C.R. Biochim. Biophys. Acta 1082:1-26(1991).
  - 18. Ribosomal protein S7e signature

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A number of eukaryotic ribosomal proteins can be grouped on the basis of sequence similarities [1]. One of these families consists of:

- Mammalian S7.
- 5 Xenopus S8.
  - Insect S7.
  - Yeast probable ribosomal protein S7 (N2212).
  - Fission yeast probable ribosomal protein S7 (SpAC18G6.13c).
- These proteins have about 200 amino acids. A highly conserved stretch of 14 residues which is located in the central section and which is rich in charged residues was selected as a signature pattern.
- 15 Consensus pattern: [KR]-L-x-R-E-L-E-K-K-F-[SAP]-x-[KR]-H
  - [1] Salazar C.E., Mills-Hamm D.M., Kumar V., Collins F.H. Nucleic Acids Res. 21:4147-4167(1993).
- 20 19. Ribosomal protein L34 signature

Ribosomal protein L34 is one of the proteins from the large subunit of the prokaryotic ribosome. It is a small basic protein of 44 to 51 amino-acid residues [1]. L34 belongs to a family of ribosomal proteins which, on the basis of sequence similarities, groups: - Eubacterial L34.

- Red algal chloroplast L34. - Cyanelle L34.

A conserved region that corresponds to the N-terminal half of L34 has been selected

as a signature pattern.

- -Consensus pattern: K-[RG]-T-[FYWL]-[EQS]-x(5)-[KRHS]-x(4,5)-G-F-x(2)-R
  - [ 1] Old I.G., Margarita D., Saint Girons I.
    Nucleic Acids Res. 20:6097-6097(1992).

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20. Ribosomal protein L6 signatures

Ribosomal protein L6 is one of the proteins from the large ribosomal subunit. In Escherichia coli, L6 is known to bind directly to the 23S rRNA and is located at the aminoacyl-tRNA binding site of the peptidyltransferase center. It belongs to a family of ribosomal proteins which, on the basis of sequence similarities [1,2,3,4], groups: - Eubacterial L6.

- Algal chloroplast L6.
- 10 Cyanelle L6.

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- Archaebacterial L6.
- Marchantia polymorpha mitochondrial L6.
- Yeast mitochondrial YmL6 (gene MRPL6).
- Mammalian L9.
- Drosophila L9.
  - Plants L9.
  - Yeast L9 (YL11).

While all the above proteins are evolutionary related it is very difficult to derive a pattern that will find them

20 all. Two patterns were therefore created, the first to detect eubacterial, cyanelle and mitochondrial L6, the second to detect archaebacterial L6 as well as eukaryotic L9.

-Consensus pattern: [PS]-[DENS]-x-Y-K-[GA]-K-G-[LIVM]

-Consensus pattern: Q-x(3)-[LIVM]-x(2)-[KR]-x(2)-R-x-F-x-D-G
[LIVM]-Y-[LIVM]-x(2)-[KR]

- [1] Suzuki K., Olvera J., Wool I.G. Gene 93:297-300(1990).
- [2] Schwank S., Harrer R., Schueller H.-J., Schweizer E. Curr. Genet. 24:136-140(1993).
- 30 [3] Golden B.L., Ramakrishnan V., White S.W. EMBO J. 12:4901-4908(1993).
  - [ 4] Otaka E., Hashimoto T., Mizuta K., Suzuki K. Protein Seq. Data Anal. 5:301-313(1993).

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21. Ribosomal protein S14p/S29e (Ribosomal protein S14 signature)

Ribosomal protein S14 is one of the proteins from the small ribosomal subunit. In Escherichia coli, S14 is known to be required for the assembly of 30S particles and may also be responsible for determining the conformation of 16S rRNA at the A site. It belongs to a family of ribosomal proteins which, on the basis of sequence similarities [1,2], groups:

- Eubacterial S14.
- Algal and plant chloroplast S14.
  - Cyanelle S14.
  - Archaebacterial Methanococcus vannielii S14.
  - Plant mitochondrial S14.
  - Yeast mitochondrial MRP2.
- Mammalian S29.

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- Yeast YS29A/B.

S14 is a protein of 53 to 115 amino-acid residues. Our signature pattern is based on the few conserved positions located in the center of these proteins.

Consensus pattern: [RP]-x(0,1)-C-x(11,12)-[LIVMF]-x-[LIVMF]-[SC]-[RG]-x(3)-[RN]

- [1] Chan Y.-L., Suzuki K., Olvera J., Wool I.G. Nucleic Acids 25 Res. 21:649-655(1993).
  - [2] Otaka E., Hashimoto T., Mizuta K. Protein Seq. Data Anal. 5:285-300(1993).
  - 22. Ribosomal protein S16 signature

Ribosomal protein S16 is one of the proteins from the small ribosomal subunit. It belongs to a family of ribosomal proteins which, on the basis of sequence similarities [1], groups:

- Eubacterial S16.

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- Algal and plant chloroplast S16.
- Cyanelle S16.

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- Neurospora crassa mitochondrial S24 (cyt-21).

S16 is a protein of about 100 amino-acid residues. A conserved region located in the N-terminal extremity of these proteins has been selected as a signature pattern.

Consensus pattern: [LIVMT]-x-[LIVM]-[KR]-L-[STAK]-R-x-G-[AKR]

10 [1] Otaka E., Hashimoto T., Mizuta K. Protein Seq. Data Anal. 5:285-300(1993).

#### 23. Ribosomal protein S21 signature

Ribosomal protein S21 is one of the proteins from the small ribosomal subunit. So far S21 has only been found in eubacteria. It is a protein of 55 to 70 amino-acid residues. A conserved region in the N-terminal section of the protein has been selected as a signature pattern.

20 Consensus pattern: [DE]-x-A-[LIY]-[KR]-R-F-K-[KR]-x(3)-[KR]

## 24. Universal stress protein family (Usp)

By a wide range of stress conditions members of the Usp family are predicted to be related to the MADS-box proteins transcript fact and bind to DNA [2]. Number of members: 39

- [1] Expression and role of the universal stress protein, UspA, of Escherichia coli during growth arrest. Nystrom T, Neidhardt FC; Mol Microbiol 1994; 11:537-544.
- 30 [2] Sequence analysis of eukaryotic developmental proteins: ancient and novel domains. Mushegian AR, Koonin EV; Genetics 1996; 144:817-828.

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#### III. Methods of Modulating Polypeptide Production

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Within the scope of invention are chimeric constructs wherein the promoter and the structural coding sequence and/or other regulatory sequences said within constructs are heterologous to each other. "Heterologous sequences" are those that are not operatively linked or are not contiguous to each other in nature. For example, a promoter from corn is considered heterologous to an Arabidopsis coding region sequence. Also, a promoter from a gene encoding a growth factor from corn is considered heterologous to a sequence encoding the corn receptor for the growth factor. Regulatory element sequences, such as UTRs or termination sequences that do not originate in nature from the the coding sequence originates from, gene as considered heterologous to said coding sequence. On the other elements operatively linked in nature are not heterologous. Thus, the promoter and coding portion of a corn gene expressing an amino acid transporter are not heterologous to each other.

Such chimeric polynucleotides are of particular interest for modulating gene expression in a host cell upon transformation of said cell with said chimeric polynucleotide.

Also within the scope of the invention are DNA molecules, whereof at least a part or portion of these DNA molecules are presented in SEQ TABLES 1 AND 2 of the present application, and wherein the structural coding sequence is under the control of its own promoter and/or its own regulatory elements. Such DNA molecules are useful for transforming the genome of a host cell or an organism regenerated from said host cell.

Typically, such polynucleotides, whether chimeric or not, are "exogenous to" the genome of an individual host cell or the organism regenerated from said host cell, such as a plant cell, respectively for a plant, when initially or subsequently introduced into said host cell or organism, by any means other

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than by a sexual cross. Examples of means by which this can be accomplished are described below, and include Agrobacteriummediated transformation (of dicots - e.g. Salomon et al. EMBO J. 3:141 (1984); Herrera-Estrella et al. EMBO J. 2:987 (1983); 5 A.C. Vergunst et al, Nucleic Acids Res. 26:11, 2729 (1998); of monocots, representative papers are those by Escudero et al., Plant J. 10:355 (1996), Ishida et al., Nature Biotechnology 14:745 (1996), May et al., Bio/Technology 13:486 (1995)), biolistic methods (Armaleo et al., Current Genetics 17:97 10 1990)), electroporation, in planta techniques, and the like. Such a plant containing the exogenous nucleic acid is referred to here as an  $R_1$  generation transgenic plant. Transgenic plants which arise from a sexual cross with another parent line or by selfing are "descendants or the progeny" of a R<sub>1</sub> plant and are 15 generally called  $F_n$  plants or  $S_n$  plants, respectively, n meaning the number of generations.

The SDFs prepared as described herein can be used to prepare expression cassettes useful in a number of techniques for suppressing or enhancing expression.

## 20 III.A. Suppression

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Expression cassettes of the invention can be used to suppress expression of endogenous genes which comprise the SDF sequence. Inhibiting expression can be useful, for instance, to tailor the ripening characteristics of a fruit (Oeller et al., Science 254:437 (1991)) or to influence seed size\_(WO98/07842) or or to provoke cell ablation (Mariani et al., Nature 357: 384-387 (1992).

As described above, a number of methods can be used to inhibit gene expression in plants, such as antisense, ribozyme, introduction of "exogenous" genes into a host cell, insertion of a polynucleotide sequence into the coding sequence and/or the promoter of the endogenous gene of interest, and the like.

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#### III.A.1. Antisense

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An expression cassette as described above can be transformed into host cell or plant to produce an antisense strand of RNA. In plant cells, it has been suggested that antisense RNA inhibits gene expression by preventing the accumulation of mRNA which encodes the enzyme of interest, see, e.g., Sheehy et al., Proc. Nat. Acad. Sci. USA, 85:8805 (1988), and Hiatt et al., U.S. Patent No. 4,801,340.

#### III.A.2. Ribozymes

Similarly, ribozyme constructs can be transformed into a plant to cleave mRNA and down-regulate translation.

#### III.A.3. Co-Suppression

Another method of suppression is by introducing an exogenous copy of the gene to be suppressed. Introduction of expression cassettes in which a nucleic acid is configured in the sense orientation with respect to the promoter has been shown to be an effective means by which to block the transcription of target genes. A detailed description of this method is described above.

# 20 III.A.4. <u>Insertion of Sequences into the Gene to</u> be Modulated

Yet another means of suppressing gene expression is to insert a polynucleotide into the gene of interest to disrupt transcription or translation of the gene.

25 Homologous recombination could<del>ean</del> be used to target a polynucleotide insert to a gene using the Cre-Lox system (A.C. Vergunst et al., *Nucleic Acids Res.* <u>26</u>:2729 (1998), A.C. Vergunst et al., *Plant Mol. Biol.* <u>38</u>:393 (1998), H. Albert et al., *Plant J.* <u>7</u>:649 (1995)).

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In addition, random insertion of polynucleotides into a host cell genome can also be used to disrupt the gene of interest. Azpiroz-Leehan et al., Trends in Genetics  $\underline{13}$ :152 (1997). In this method, screening for clones from a library containing random insertions is preferred for identifying those that have polynucleotides inserted into the gene of interest. Such screening can be performed using probes and/or primers described above based on sequences from SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequence thereto. The screening can also be performed by selecting clones or  $R_1$  plants having a desired phenotype.

## III.A.5. Promoter Modulation

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Inactivation of the promoter that drives a gene of interest can modulate transcription and translation, and therefore expression. For example, triple helices can be formed using oligonucleotides based on sequences from SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequence thereto. The oligonucleotide can be delivered to the host cell can bind to the promoter in the genome to form a triple helix and prevent transcription.

Additionally, a vector capable of producing the oligonucleotide can be inserted into the host cell to deliver the oligonucleotide.

#### III.A.6. Expression of Mutants

An alternative method for inhibiting gene function is through the use of dominant negative mutations. Dominant negative mutations produce a mutant polypeptide which is capable of competing with the native polypeptide, but which does not produce the native result. Consequently, over expression of these mutations can titrate out an undesired activity of the native protein. For example, the inactive dominant-negative mutant may bind to the same receptor as the

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native protein, preventing the native protein from activating a signal transduction pathway. Alternatively, the dominant-negative mutant can be an inactive enzyme still capable of binding to the same substrate as the native protein.

Dominant-negative mutants also can act upon the native protein itself to prevent activity. For example, the native protein may be active only as a homo-multimer or as one subunit of a hetero-multimer. Incorporation of an inactive subunit into the multimer with native subunit(s) can inhibit activity.

Thus, gene function can be modulated by insertion of an expression construct encoding a dominant-negative mutant into a host cell of interest.

#### III.B. Enhanced Expression

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Enhanced expression of a gene of interest in a host cell can be accomplished by either (1) insertion of an exogenous gene; or (2) promoter modulation.

#### III.B.1. Insertion of an Exogenous Gene

Insertion of an expression construct encoding an exogenous gene can boost the number of gene copies expressed in a host cell.

Such expression constructs can comprise genes that either encode the native protein that is of interest or that encode a variant that exhibits enhanced activity as compared to the native protein. Such genes encoding proteins of interest can be constructed from the sequences from SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequence thereto.

Such an exogenous gene can include either a constitutive promoter permitting expression in any cell in a host organism or a promoter that directs expression only in particular cells or times during a host cell life cycle or in response to environmental stimuli.

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## III.B.2. Promoter Modulation

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Some promoters require binding of a regulatory protein to be activated. Other promoters may need a protein that signals a promoter binding protein to expose a polymerase binding site. In either case, over-expression of such proteins can be used to enhance expression of a gene of interest by increasing the activation time of the promoter.

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Such regulatory proteins are encoded by some of the sequences in SEQ TABLES 1 AND 2, fragments thereof, and substantially similar sequences thereto.

Coding sequences for these proteins can be constructed as described above.

In some cases, duplication of enhancer elements or insertion of exogenous enhancer elements will increase expression of a desired gene from a particular promoter. The useful enhancer elements can be portions of one or more of the SDFs of SEQ TABLES 1 AND 2.

#### IV. Gene Constructs and Vector Construction

To use isolated SDFs of the present invention or a combination of them or parts and/or mutants and/or fusions of said SDFs in the above techniques, recombinant DNA vectors which comprise said SDFs and are suitable for transformation of cells, such as plant cells, are usually prepared.

The vector backbone can be any of those typical in the art such as plasmids, viruses, artificial chromosomes, BACs, YACs and PACs and vectors of the sort described by \*\*.

Typically, a vector will comprise the exogenous gene, which in its turn comprises an SDF of the present invention to be introduced into the genome of a host cell, and which gene may be an antisense construct, a ribozyme construct, or a structural coding sequence with any desired transcriptional and/or translational regulatory sequences, such as promoters

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and 3' end termination sequences. Vectors of the invention can also include origins of replication, markers, homologous

sequences, introns, etc.

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A DNA sequence coding for the desired polypeptide, for example a cDNA sequence encoding a full length protein, will preferably be combined with transcriptional and translational initiation regulatory sequences which will direct the transcription of the sequence from the gene in the intended tissues of the transformed plant.

For example, for over-expression, a plant promoter fragment may be employed that will direct expression of the gene in all tissues of a regenerated plant. Such promoters are referred to herein as "constitutive" promoters and are active under most environmental conditions and states of development or cell differentiation. Examples of constitutive promoters include the cauliflower mosaic virus (CaMV) 35S transcription initiation region, the 1' or 2' promoter derived from T-DNA of Agrobacterium tumefaciens, and other transcription initiation regions from various plant genes known to those of skill.

Alternatively, the plant promoter may direct expression of an SDF of the invention in а specific (tissue-specific promoters) or may be otherwise under more precise environmental control (inducible promoters). Examples tissue-specific promoters under developmental control include promoters that initiate transcription only in certain tissues, such as root, ovule, fruit, seeds, or flowers. promoter from a LEC1 gene, described in copending application U.S. Ser. No. 09/103,478, is particularly useful for directing gene expression so that a desired gene product is located in embryos or seeds. Other suitable promoters include those from genes encoding storage proteins or the lipid body membrane protein, oleosin. A few root-specific promoters are noted above. Examples of environmental conditions that may affect

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transcription by inducible promoters include anaerobic conditions, elevated temperature, or the presence of light.

If proper polypeptide expression is desired, a polyadenylation region at the 3'-end of the coding region should be included. The polyadenylation region can be derived from the natural gene, from a variety of other plant genes, or from T-DNA.

The vector comprising the sequences (e.g., promoters or coding regions) from genes of the invention will typically comprise a marker gene that confers a selectable phenotype on plant cells. For example, the marker may encode biocide resistance, particularly antibiotic resistance, such as resistance to kanamycin, G418, bleomycin, hygromycin, or herbicide resistance, such as resistance to chlorosulfuron or phosphinotricin.

#### IV.A.Coding Sequences

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Generally, the sequence in the transformation vector and to be introduced into the genome of the host cell does not need to be absolutely identical to an SDF of the present invention. Also, it is not necessary for it to be full length, relative to either the primary transcription product or fully processed mRNA. Use of sequences shorter than full-length may be preferred to avoid concurrent production of some plants that are overexpressors. Furthermore, the introduced sequence need not have the same intron or exon pattern as a native gene. Also, heterologous non-coding segments can be incorporated into the coding sequence without changing the desired amino acid sequence of the polypeptide to be produced.

#### IV.B. Promoters

As explained above, introducing an exogenous SDF from the same species or an orthologous SDF from another species can modulate the expression of a native gene corresponding to

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that SDF of interest. Such an SDF construct can be under the control of either a constitutive promoter (e.g., the promoter of the 35S gene of the cauliflower mosaic virus or the promotor of the gene encoding the cowpea trypsin inhibitor) or a highly regulated inducible promoter (e.g., a copper inducible promoter). The promoter of interest can initially be either endogenous or heterologous to the species question. When re-introduced into the genome of said species, such promoter becomes "exogenous" to said species. The promoter-SDF construct can be made using standard recombinant DNA techniques (Sambrook et al. 1989) and can be introduced the species of interest by Agrobacterium-mediated to transformation or by other means of transformation (e.g., particle qun bombardment) as referenced above. expression of an SDF transgene can lead to co-suppression of the homologous gene thereby creating some alterations in the phenotypes of the transformed species as demonstrated by similar analysis of the chalcone synthase gene (Napoli et al., Plant Cell 2:279 (1990) and van der Krol et al., Plant Cell 2:291 (1990)). If an SDF is found to encode a protein with desirable characteristics, its over-expression can be controlled so that its accumulation can be manipulated in an organ- or tissue-specific manner utilizing a promoter having such specificity.

Likewise, if the promoter of an SDF (or an SDF that includes a promoter) is found to be tissue-specific or developmentally regulated, such a promoter can be utilized to drive the expression of a specific gene of interest (e.g., seed storage protein or root-specific protein). Thus, the level of accumulation of a particular protein can be manipulated or its spatial localization in an organ- or tissue-specific manner can be altered.

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## IV. C Signal Peptides

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In some cases it may be desirable for the protein encoded by an introduced exogenous or orthologous SDF to be targeted (1) to a particular organelle, (2) to interact with a particular molecule or (3) for secretion outside of the cell harboring the introduced SDF. This will be accomplished using a signal peptide.

Signal peptides direct protein targeting, are involved in ligand-receptor interactions and act in cell to cell communication. Many proteins, especially soluble proteins, contain a signal peptide that targets the protein to one of several different intracellular compartments. In plants, these compartments include, but are not limited to, the endoplasmic reticulum (ER), mitochondria, plastids (such as chloroplasts), the vacuole, the Golgi apparatus, protein storage vessicles (PSV) and, in general, membranes . signal peptide sequences are conserved, such as the Asn-Pro-Ile-Arg amino acid motif found in the N-terminal propeptide signal that targets proteins to the vacuole (Marty (1999) The Plant Cell 11: 587-599). Other signal peptides do not have a consensus sequence per se, but are largely composed of hydrophobic amino acids, such as those signal peptides targeting proteins to the ER (Vitale and Denecke (1999) The Plant Cell 11: 615-628). Still others do not appear to contain either a consensus sequence or an identified common secondary sequence, for instance the chloroplast stromal targeting signal peptides (Keegstra and Cline (1999) The Plant Cell 11: 557-570). Furthermore, some targeting peptides are bipartite, directing proteins first to an organelle and then to a membrane within the organelle (e.g. within the thylakoid lumen of the chloroplast; see Keegstra and Cline (1999) The Plant Cell 11: 557-570). In addition to the diversity in sequence and secondary structure, placement of the signal peptide is also varied. Proteins destined for

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the vacuole, for example, have targeting signal peptides found at the N-terminus, at the C-terminus and at a surface location in mature, folded proteins.

Signal peptides also serve as ligands for some receptors. Perhaps the best known example of this is the interaction of the ER targeting signal peptide with the signal recognition particle (SRP). Here, the SRP binds to the signal peptide, halting translation, and the resulting SRP complex then binds to docking proteins located on the surface of the ER, prompting the transfer of the protein into the ER.

These characteristics of signal proteins can be used to more tightly control the expression of introduced SDFs. particular, associating the appropriate signal sequence with a specific SDF can allow sequestering of the protein in specific organelles (plastids, as an example), secretion outside of the cell, targeting interaction with particular receptors, etc. Hence, the inclusion of signal proteins in constructs involving the SDFs of the invention increases the range of manipulation of SDF expression. To carry this out, constructs are made with the nucleotide sequence of a known signal peptide immediately 5' to the initiation of the coding region of an SDF so that the signal peptide is translated in frame with the coding region and immediately precedes it. The nucleotide sequence of the signal peptide can be isolated from characterized genes using common molecular biological techniques or can be synthesized in vitro.

#### V. Transformation Techniques

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A wide range of techniques for inserting exogenous 30 polynucleotides are known for a number of host cells, including, without limitation, bacterial, yeast, mammalian, insect and plant cells.

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Techniques for transforming a wide variety of higher plant species are well known and described in the technical and scientific literature. See, e.g. Weising et al., Ann. Rev. Genet. 22:421 (1988); and Christou, Euphytica, v. 85, n.1-3:13-27, (1995).

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DNA constructs of the invention may be introduced into genome of the desired plant host by a variety of conventional techniques. For example, the DNA construct may be introduced directly into the genomic DNA of the plant cell using techniques such as electroporation and microinjection of plant cell protoplasts, or the DNA constructs can be introduced directly to plant tissue using ballistic methods, such as DNA particle bombardment. Alternatively, the DNA constructs may be combined with suitable T-DNA flanking regions and introduced into a conventional Agrobacterium tumefaciens host vector. The virulence functions of the Agrobacterium tumefaciens host will direct the insertion of the construct and adjacent marker into the plant cell DNA when the cell is infected by the bacteria (Vergunst et al., Nucl. Acids. Res. 26:2729 (1998) (sitedirected integration using a Cre-Lox recombinase system); McCormac et al., Mol. Biotechnol. 8:199 (1997); Hamilton, Gene 200:107 (1997)); Salomon et al. EMBO J. 3:141 (1984); Herrera-Estrella et al. EMBO J. 2:987 (1983).

Microinjection techniques are known in the art and well the scientific and patent literature. introduction of DNA constructs using polyethylene glycol precipitation is described in Paszkowski et al. EMBO J. 3:2717 Electroporation techniques are described in Fromm et Acad. Sci. USA 82:5824 (1985). al. Proc. Natl transformation techniques are described in Klein et al. Nature tumefaciens-mediated 327:773 (1987).Agrobacterium transformation techniques, including disarming and use of binary vectors, well described in the scientific are literature. See, for example Hamilton, CM., Gene 200:107

(1997); Müller et al. *Mol. Gen. Genet.* 207:171 (1987); Komari et al. *Plant J.* 10:165 (1996); Venkateswarlu et al. *Biotechnology* 9:1103 (1991) and Gleave, *AP., Plant Mol. Biol.* 20:1203 (1992); Graves and Goldman, *Plant Mol. Biol.* 7:34 (1986) and Gould et al., *Plant Physiology* 95:426 (1991).

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Transformed plant cells which are derived by any of the above transformation techniques can be cultured to regenerate a whole plant that possesses the transformed genotype and thus the desired phenotype such as seedlessness. Such regeneration techniques rely on manipulation of certain phytohormones in a tissue culture growth medium, typically relying on a biocide and/or herbicide marker which has been introduced together with the desired nucleotide sequences. Plant regeneration cultured protoplasts is described in Evans et al., Protoplasts Isolation and Culture in "Handbook of Plant Cell Culture," pp. 124-176, MacMillan Publishing Company, New York, 1983; and Binding, Regeneration of Plants, Plant Protoplasts, pp. 21-73, CRC Press, Boca Raton, 1988. Regeneration can also be obtained from plant callus, explants, organs, or parts thereof. regeneration techniques are described generally in Klee et al. Ann. Rev. of Plant Phys. 38:467 (1987). Regeneration of monocots (rice) is described by Hosoyama et al. (Biosci. Biotechnol. Biochem. 58:1500 (1994)) and by Ghosh et al. (J. Biotechnol. 32:1 (1994)). The nucleic acids of the invention can be used to confer desired traits on essentially any plant.

Thus, the invention has use over a broad range of plants, including species from the genera Asparagus, Atropa, Avena, Brassica, Citrus, Citrullus, Capsicum, Cucumis, Cucurbita, Daucus, Fragaria, Glycine, Gossypium, Helianthus, Heterocallis, Hordeum, Hyoscyamus, Lactuca, Linum, Lolium, Lycopersicon, Malus, Manihot, Majorana, Medicago, Nicotiana, Oryza, Panieum, Pannesetum, Persea, Pisum, Pyrus, Prunus, Raphanus, Secale, Senecio, Sinapis, Solanum, Sorghum, Trigonella, Triticum, V'tis, Vigna, and, Zea.

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One of skill will recognize that after the expression cassette is stably incorporated in transgenic plants and confirmed to be operable, it can be introduced into other plants by sexual crossing. Any of a number of standard breeding techniques can be used, depending upon the species to be crossed.

The particular sequences of SDFs identified are provided in the attached SEQ TABLES 1 AND 2. One of ordinary skill in the art, having this data, can obtain cloned DNA fragments, synthetic DNA fragments or polypeptides constituting desired sequences by recombinant methodology known in the art or described herein.

#### **EXAMPLES**

The invention is illustrated by way of the following examples. The invention is not limited by these examples as the scope of the invention is defined solely by the claims following.

#### EXAMPLE 1: SOUTHERN HYBRIDIZATIONS

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The SDFs of the invention can be used in Southern hybridizations as described above. The following describes extraction of DNA from nuclei of plant cells, digestion of the nuclear DNA and separation by length, transfer of the separated fragments to membranes, preparation of probes for hybridization, hybridization and detection of the hybridized probe.

The procedures described herein can be used to isolate related polynucleotides or for diagnostic purposes. Moderate stringency hybridization conditions, as defined above, are described in the present example. These conditions result in detection of hybridization between sequences having at least 70% sequence identity. As described above, the hybridization and wash conditions can be changed to reflect the desired

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degree of sequence identity between probe and target sequences that can be detected.

In the following procedure, a probe for the hybridization is produced from two PCR reactions using two primers from genomic sequence of *Arabidopsis thaliana*. As described above, the particular template for generating the probe can be any desired template.

The first PCR product is assessed to validate the size of the primer to assure it is of the expected size. Then the product of the first PCR is used as a template, with the same pair of primers used in the first PCR, in a second PCR that produces a labeled product used as the probe.

Fragments detected by hybridization, or other bands of interest, can be isolated from gels used to separate genomic DNA fragments by known methods for further purification and/or characterization.

#### Buffers for nuclear DNA extraction

## 1. 10X HB

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	1000 ml	
40 mM spermidine	10.2 g	Spermine (Sigma S-2876) and spermidine (Sigma S-2501)
10 mM spermine	3.5 g	Stabilize chromatin and the nuclear membrane
0.1 M EDTA (disodium)	37.2 g	EDTA inhibits nuclease
0.1 M Tris	12.1 g	Buffer
0.8 M KCl	59.6 g	Adjusts ionic strength for stability of nuclei

Adjust pH to 9.5 with 10 N NaOH. It appears that there is a nuclease present in leaves. Use of pH 9.5 appears to inactivate this nuclease.

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2. 2 M sucrose (684 g per 1000 ml)

Heat about half the final volume of water to about 50°C.

Add the sucrose slowly then bring the mixture to close to final volume; stir constantly until it has dissolved.

Bring the solution to volume.

3. Sarkosyl solution (lyses nuclear membranes)

1000 ml

N-lauroyl sarcosine (Sarkosyl) 20.0 g

0.1 M Tris 12.1 g

10 0.04 M EDTA (Disodium) 14.9 q

Adjust the pH to 9.5 after all the components are dissolved and bring up to the proper volume.

4. 20% Triton X-100 80 ml Triton X-100

15 320 ml 1xHB (w/o  $\beta$ -ME and PMSF) Prepare in advance; Triton takes some time to dissolve

A. Procedure

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1. Prepare 1X "H" buffer (keep ice-cold during use)

20 10X HB 100 ml
2 M sucrose 250 ml a non-ionic osmoticum
Water 634 ml

#### Added just before use:

100 mM PMSF\*

10 ml a protease
inhibitor; protects
nuclear membrane proteins

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ß-mercaptoethanol 1 ml inactivates nuclease by reducing disulfide bonds

\*100 mM PMSF

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(phenyl methyl sulfonyl fluoride, Sigma P-7626) (add 0.0875 g to 5 ml 100% ethanol)

- 2. Homogenize the tissue in a blender (use 300-400 ml of 1xHB per blender). Be sure that you use 5-10 ml of HB buffer per gram of tissue. Blenders generate heat so be sure to keep the homogenate cold. It is necessary to put the blenders in ice periodically.
  - 3. Add the 20% Triton X-100 (25 ml per liter of homogenate) and gently stir on ice for 20 min. This lyses plastid, but not nuclear, membranes.
- 15 4. Filter the tissue suspension through several nylon filters into an ice-cold beaker. The first filtration is through a 250-micron membrane; the second is through an 85-micron membrane; the third is through a 50-micron membrane; and the fourth is through a 20-micron membrane. Use a large funnel to hold the filters. Filtration can be sped up by gently squeezing the liquid through the filters.
  - 5. Centrifuge the filtrate at  $1200 \times g$  for 20 min. at  $4^{\circ}\text{C}$  to pellet the nuclei.
- 25 6. Discard the dark green supernatant. The pellet will have several layers to it. One is starch; it is white and gritty. The nuclei are gray and soft. In the early

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steps, there may be a dark green and somewhat viscous layer of chloroplasts.

Wash the pellets in about 25 ml cold H buffer (with Triton X-100) and resuspend by swirling gently and pipetting. After the pellets are resuspended.

Pellet the nuclei again at  $1200 - 1300 \times g$ . Discard the supernatant.

Repeat the wash 3-4 times until the supernatant has changed from a dark green to a pale green. This usually happens after 3 or 4 resuspensions. At this point, the pellet should be grayish white and very slippery. The Triton X-100 in these repeated steps helps to destroy the chloroplasts and mitochondria that contaminate the prep.

- Resuspend the nuclei for a final time in a total of 15 ml of H buffer and transfer the suspension to a sterile 125 ml Erlenmeyer flask.
- 7. Add 15 ml, dropwise, cold 2% Sarkosyl, 0.1 M Tris, 0.04 M EDTA solution (pH 9.5) while swirling gently. This lyses the nuclei. The solution will become very viscous.
  - 8. Add 30 grams of CsCl and gently swirl at room temperature until the CsCl is in solution. The mixture will be gray, white and viscous.
- 25 9. Centrifuge the solution at  $11,400 \times g$  at  $4^{\circ}C$  for at least 30 min. The longer this spin is, the firmer the protein pellicle.

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- The result should be a clear green supernatant over a 10. white pellet, and (perhaps) under a protein pellicle. Carefully remove the solution under the protein pellicle and above the pellet. Determine the density of the solution by weighing 1 ml of solution and add CsCl if necessary to bring to 1.57 g/ml. The solution contains dissolved solids (sucrose etc) and the refractive index alone will not be an accurate quide to CsCl concentration.
  - 11. Add 20  $\mu$ l of 10 mg/ml EtBr per ml of solution.
  - 12. Centrifuge at  $184,000 \times g$  for 16 to 20 hours in a fixed-angle rotor.
- 13. Remove the dark red supernatant that is at the top of the tube with a plastic transfer pipette and discard. Carefully remove the DNA band with another transfer pipette. The DNA band should be visible in room light; otherwise, use a long wave UV light to locate the band.
- 14. Extract the ethidium bromide with isopropanol saturated with water and salt. Once the solution is clear, extract at least two more times to ensure that all of the EtBr is gone. Be very gentle, as it is very easy to shear the DNA at this step. This extraction may take a while because the DNA solution tends to be very viscous.

  25 If the solution is too viscous, dilute it with TE.
  - 15. Dialyze the DNA for at least two days against several changes (at least three times) of TE (10 mM Tris, 1mM EDTA, pH 8) to remove the cesium chloride.

16. Remove the dialyzed DNA from the tubing. If the dialyzed DNA solution contains a lot of debris, centrifuge the DNA solution at least at 2500 x g for 10 min. and carefully transfer the clear supernatant to a new tube. Read the A260 concentration of the DNA.

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17. Assess the quality of the DNA by agarose gel electophesis (1% agarose gel) of the DNA. Load 50 ng and 100 ng (based on the OD reading) and compare it with known and good quality DNA. Undigested lambda DNA and a lambda-HindIII-digested DNA are good molecular weight makers.

## Protocol for Digestion of Genomic DNA

## Protocol:

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- 1. The relative amounts of DNA for different crop plants that provide approximately a balanced number of genome equivalent is given in Table 3. Note that due to the size of the wheat genome, wheat DNA will be underrepresented. Lambda DNA provides a useful control for complete digestion.
- 2. Precipitate the DNA by adding 3 volumes of 100% ethanol.

  20 Incubate at -20°C for at least two hours. Yeast DNA can be purchased and made up at the necessary concentration, therefore no precipitation is necessary for yeast DNA.
- 3. Centrifuge the solution at 11,400 x g for 20 min.

  Decant the ethanol carefully (be careful not to disturb
  the pellet). Be sure that the residual ethanol is
  completely removed either by vacuum desiccation or by
  carefully wiping the sides of the tubes with a clean
  tissue.

4. Resuspend the pellet in an appropriate volume of water.

Be sure the pellet is fully resuspended before proceeding to the next step. This may take about 30

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min.

- 5 5. Add the appropriate volume of 10X reaction buffer provided by the manufacturer of the restrictioned-enzyme to the resuspended DNA followed by the appropriate volume of enzymes. Be sure to mix it properly by slowly swirling the tubes.
- 10 6. Set-up the lambda digestion-control for each DNA that you are digesting.
  - 7. Incubate both the experimental and lambda digests overnight at  $37\,^{\circ}\text{C}$ . Spin down condensation in a microfuge before proceeding.
- 15 8. After digestion, add 2  $\mu$ l of loading dye (typically 0.25% bromophenol blue, 0.25% xylene cyanol in 15% Ficoll or 30% glycerol) to the lambda-control digests and load in 1% TPE-agarose gel (TPE is 90 mM Trisphosphate, 2 mM EDTA, pH 8). If the lambda DNA in the lambda control digests are completely digested, proceed with the precipitation of the genomic DNA in the digests.
- 9. Precipitate the digested DNA by adding 3 volumes of 100% ethanol and incubating in -20%C for at least 2 hours (preferably overnight).
  - EXCEPTION: Arabidopsis and yeast DNA are digested in an appropriate volume; they don't have to be precipitated.

10. Resuspend the DNA in an appropriate volume of TE (e.g., 22  $\mu$ l x 50 blots = 1100  $\mu$ l) and an appropriate volume of 10X loading dye (e.g., 2.4  $\mu$ l x 50 blots = 120  $\mu$ l). Be careful in pipetting the loading dye - it is viscous. Be sure you are pipetting the correct volume.

Table 3
Some guide points in digesting genomic DNA.

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Species	Genome Size	Size Relative to Arabidopsis	Genome Equivalent to 2 µg Arabidopsis DNA	Amount of DNA per blot
Arabidopsis	120 Mb	1X	1X	2 μg
Brassica	1,100 Mb	9.2X	0.54X	10 μg
Corn	2,800 Mb	23.3X	0.43X	20 μg
Cotton	2,300 Mb	19.2X	0.52X	20 μg
Oat	11,300 Mb	94X	0.11X	20 µg
Rice	400 Mb	3.3X	0.75X	5 μg
Soybean	1,100 Mb	9.2X	0.54X	10 μg
Sugarbeet	758 Mb	6.3X	0.8X	10 μg
Sweetclover	1,100 Mb	9.2X	0.54X	10 μg
Wheat	16,000 Mb	133X	0.08X	20 μg
Yeast	15 Mb	0.12X	1X	0.25 μg

## 10 Protocol for Southern Blot Analysis

The digested DNA samples are electrophoresed in 1% agarose gels in 1x TPE buffer. Low voltage; overnight separations are preferred. The gels are stained with EtBr and photographed.

1. For blotting the gels, first incubate the gel in  $0.25\ N$  HCl (with gentle shaking) for about 15 min.

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- Then briefly rinse with water. The DNA is denatured by 2 incubations. Incubate (with shaking) in 0.5 M NaOH in 1.5 M NaCl for 15 min.
- 3. The gel is then briefly rinsed in water and neutralized by incubating twice (with shaking) in 1.5 M Tris pH 7.5 in 1.5 M NaCl for 15 min.
  - 4. A nylon membrane is prepared by soaking it in water for at least 5 min, then in 6X SSC for at least 15 min. before use. (20x SSC is 175.3 g NaCl, 88.2 g sodium citrate per liter, adjusted to pH 7.0.)
  - 5. The nylon membrane is placed on top of the gel and all bubbles in between are removed. The DNA is blotted from the gel to the membrane using an absorbent medium, such as paper toweling and 6x SCC buffer. After the transfer, the membrane may be lightly brushed with a gloved hand to remove any agarose sticking to the surface.
- 6. The DNA is then fixed to the membrane by UV crosslinking and baking at  $80^{\circ}\text{C}$ . The membrane is stored at  $4^{\circ}\text{C}$  until use.
  - B. Protocol for PCR Amplification of Genomic Fragments in Arabidopsis

#### Amplification procedures:

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1. Mix the following in a 0.20 ml PCR tube or 96-well PCR plate:

		Final Amount or
Volume	Stock	Conc.

0.5 μ1	~ 10 ng/ $\mu$ l genomic DNA <sup>1</sup>	5 ng
2.5 μl	10X PCR buffer	20 mM Tris, 50 mM KCl
0.75 μl	50 mM MgCl <sub>2</sub>	1.5 mM
1 μ1	10 pmol/µl Primer 1 (Forward)	10 pmol
1 μ1	10 pmol/µl Primer 2 (Reverse)	10 pmol
0.5 μl	5 mM dNTPs	0.1 mM
0.1 μ1	5 units/µl Platinum Taq™ (Life Technologies, Gaithersburg, MD) DNA Polymerase	1 units
(to 25 μ1)	Water	

- 2. The template DNA is amplified using a Perkin Elmer 9700 PCR machine.
- 1) 94°C for 10 min. followed by

2)	3)	4)
5 cycles:	5 cycles:	25 cycles:
94 °C - 30 sec	94 °C - 30 sec	94 °C - 30 sec
62°C - 30 sec	58°C - 30 sec	53 °C - 30 sec
72°C - 3 min	72°C - 3 min	72°C - 3 min

5)  $72^{\circ}\text{C}$  for 7 min. Then the reactions are stopped by chilling to  $4^{\circ}\text{C}$ .

<sup>&</sup>lt;sup>1</sup> Arabidopsis DNA is used in the present experiment, but the procedure is a general one.

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The procedure can be adapted to a multi-well format if necessary.

#### Quantification and Dilution of PCR Products:

- 1. The product of the PCR is analyzed by electrophoresis in a 1% agarose gel. A linearized plasmid DNA can be used as a quantification standard (usually at 50, 100, 200, and 400 ng). These will be used as references to approximate the amount of PCR products. HindIII-digested Lambda DNA is useful as a molecular weight marker. The gel can be run fairly quickly; e.g., at 100 volts. The standard gel is examined to determine that the size of the PCR products is consistent with the expected size and if there are significant extra bands or smeary products in the PCR reactions.
- 15 2. The amounts of PCR products can be estimated on the basis of the plasmid standard.
  - 3. For the small number of reactions that produce extraneous bands, a small amount of DNA from bands with the correct size can be isolated by dipping a sterile  $10-\mu l$  tip into the band while viewing though a UV Transilluminator. The small amount of agarose gel (with the DNA fragment) is used in the labeling reaction.

#### C. Protocol for PCR-DIG-Labeling of DNA

#### Solutions:

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Reagents in PCR reactions (diluted PCR products, 10X PCR Buffer, 50 mM MgCl $_2$ , 5 U/ $\mu$ l Platinum Taq Polymerase, and the primers)

10X dNTP + DIG-11-dUTP [1:5]: (2 mM dATP, 2 mM dCTP, 2 mM dGTP, 1.65 mM dTTP, 0.35 mM DIG-11-dUTP)

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10X dNTP + DIG-11-dUTP [1:10]: (2 mM dATP, 2 mM dCTP, 2 mM dGTP, 1.81 mM dTTP, 0.19 mM DIG-11-dUTP)

10X dNTP + DIG-11-dUTP [1:15]: (2 mM dATP, 2 mM dCTP, 2 mM dGTP, 1.875 mM dTTP, 0.125 mM DIG-11-dUTP)

5 TE buffer (10 mM Tris, 1 mM EDTA, pH 8)

Maleate buffer: In 700 ml of deionized distilled water, dissolve 11.61 g maleic acid and 8.77 g NaCl. Add NaOH to adjust the pH to 7.5. Bring the volume to 1 L. Stir for 15 min. and sterilize.

10% blocking solution: In 80 ml deionized distilled water, dissolve 1.16g maleic acid. Next, add NaOH to adjust the pH to 7.5. Add 10 g of the blocking reagent powder (Boehringer Mannheim, Indianapolis, IN, Cat. no. 1096176). Heat to 60°C while stirring to dissolve the powder. Adjust the volume to 100 ml with water. Stir and sterilize.

1% blocking solution: Dilute the 10% stock to 1% using the maleate buffer.

Buffer 3 (100 mM Tris, 100 mM NaCl, 50 mM MgCl<sub>2</sub>, pH9.5).

20 Prepared from autoclaved solutions of 1M Tris pH 9.5, 5

M NaCl, and 1 M MgCl<sub>2</sub> in autoclaved distilled water.

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#### Procedure:

1. PCR reactions are performed in 25  $\mu$ l volumes containing:

PCR buffer 1X

 $MgCl_2$  1.5 mM

5 10X dNTP + DIG-11-dUTP 1X (please see the

note below)

Platinum Taq<sup>™</sup> Polymerase 1 unit

10 pg probe DNA

10 pmol primer 1

Note: <u>Use for</u>:

10X dNTP + DIG-11-dUTP (1:5) < 1 kb 10X dNTP + DIG-11-dUTP (1:10) 1 kb to 1.8 kb

10X dNTP + DIG-11-dUTP (1:15) > 1.8 kb

- 2. The PCR reaction uses the following amplification cycles:
  - 1) 94°C for 10 min.

2)	3)	4)
5 cycles:	5 cycles:	25 cycles:
95°C - 30 sec	95°C - 30 sec	95°C - 30 sec
61°C - 1 min	59°C - 1 min	51°C - 1 min
73°C - 5 min	75°C - 5 min	73°C - 5 min

- 5)  $72^{\circ}\text{C}$  for 8 min. The reactions are terminated by chilling to  $4^{\circ}\text{C}$  (hold).
- 3. The products are analyzed by electrophoresis- in a 1% agarose gel, comparing to an aliquot of the unlabelled probe starting material.

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4. The amount of DIG-labeled probe is determined as follows:

Make serial dilutions of the diluted control DNA in dilution buffer (TE: 10 mM Tris and 1 mM EDTA, pH 8) as shown in the following table:

DIG-labeled control DNA starting conc.	Stepwise Dilution	Final Conc. (Dilution Name)
5 ng/µl	1 μl in 49 μl TE	100 pg/μl (A)
100 pg/μl (A)	25 µl in 25 µl TE	50 pg/μl (B)
50 pg/µl (B)	25 µl in 25 µl TE	25 pg/μl (C)
25 pg/μl (C)	20 μl in 30 μl TE	10 pg/μl (D)

- a. Serial deletions of a DIG-labeled standard DNA ranging from 100 pg to 10 pg are spotted onto a positively charged nylon membrane, marking the membrane lightly with a pencil to identify each dilution.
- b. Serial dilutions (e.g., 1:50, 1:2500, 1:10,000) of the newly labeled DNA probe are spotted.
- c. The membrane is fixed by UV crosslinking.
- 15 d. The membrane is wetted with a small amount of maleate buffer and then incubated in 1% blocking solution for 15 min at room temp.
  - e. The labeled DNA is then detected using alkaline phosphatase conjugated anti-DIG antibody (Boehringer Mannheim, Indianapolis, IN, cat. no.

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1093274) and an NBT substrate according to the manufacture's instruction.

f. Spot intensities of the control and experimental dilutions are then compared to estimate the concentration of the PCR-DIG-labeled probe.

## D. Prehybridization and Hybridization of Southern Blots Solutions:

100% Formamide purchased from Gibco

20X SSC (1X = 0.15 M NaCl, 0.015 M

 $Na_3$ citrate)

per L: 175 g NaCl

87.5 g Na<sub>3</sub>citrate·2H<sub>2</sub>0

20% Sarkosyl (N-lauroyl-sarcosine)

20% SDS (sodium dodecyl sulphate)

10% Blocking Reagent: In 80 ml deionized distilled water, dissolve 1.16 g maleic acid. Next, add NaOH to adjust the pH to 7.5. Add 10 g of the blocking reagent powder. Heat to 60°C while stirring to dissolve the powder. Adjust the volume to 100 ml with water. Stir and sterilize.

Prehybridization Mix:

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Final		Volume	
Concentration	Components	(per 100 ml)	Stock
50%	Formamide	50 ml	100%
5X	SSC	25 ml	20X
0.1%	Sarkosyl	0.5 ml	20%

-	_	
7	$\alpha$	
- 1		

0.02%	SDS	0.1 ml	20%
2%	Blocking Reagent	20 ml	10%
	Water	4.4 ml	

#### General Procedures:

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- Place the blot in a heat-sealable plastic bag and add an 1. appropriate volume of prehybridization solution ml/100cm<sup>2</sup>) at room temperature. Seal the bag with a heat sealer, avoiding bubbles as much as possible. Lav down the bags in large plastic tray (one tray a can accommodate at least 4-5 bags). Ensure that the bags are lying flat in the tray so that the prehybridization evenly distributed throughout the bag. solution is Incubate the blot for at least 2 hours with gentle agitation using a waver shaker.
- 2. Denature DIG-labeled DNA probe by incubating for 10 min. at  $98\,^{\circ}\text{C}$  using the PCR machine and immediately cool it to  $4\,^{\circ}\text{C}$ .
- 3. Add probe to prehybridization solution (25 ng/ml; 30 ml = 750 ng total probe) and mix well but avoid foaming. Bubbles may lead to background.
- 4. Pour off the prehybridization solution from the hybridization bags and add new prehybridization and probe solution mixture to the bags containing the membrane.
  - 5. Incubate with gentle agitation for at least 16 hours.
  - 6. Proceed to medium stringency post-hybridization wash:

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Three times for 20 min. each with gentle agitation using  $1 \times SSC$ ,  $1 \times SDS$  at  $60 \circ C$ .

All wash solutions must be prewarmed to 60°C. Use about 100 ml of wash solution per membrane.

To avoid background keep the membranes fully submerged to avoid drying in spots; agitate sufficiently to avoid having membranes stick to one another.

7. After the wash, proceed to immunological detection and CSPD development.

# 10 E. Procedure for Immunological Detection with CSPD Solutions:

Buffer 1: Maleic acid buffer (0.1 M maleic

acid, 0.15 M NaCl; adjusted to pH

7.5 with NaoH)

15 Washing buffer: Maleic acid buffer with 0.3% (v/v)

Tween 20.

Dissolve

Blocking stock solution 10% blocking reagent in buffer 1.

blocking reagent powder (Boehringer Mannheim, Indianapolis, IN, cat. no.

concentration):

(10X

1096176) by constantly stirring on a 65°C heating block or heat in a microwave, autoclave and store at

4°C.

25 Buffer 2

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(1X blocking solution): Dilute the stock solution 1:10 in Buffer 1.

Detection buffer: 0.1 M Tris, 0.1 M NaCl, pH 9.5

#### Procedure:

- 5 1. After the post-hybridization wash the blots are briefly rinsed (1-5 min.) in the maleate washing buffer with gentle shaking.
  - 2. Then the membranes are incubated for 30 min. in Buffer 2 with gentle shaking.
- 10 3. Anti-DIG-AP conjugate (Boehringer Mannheim, Indianapolis, IN, cat. no. 1093274) at 75 mU/ml (1:10,000) in Buffer 2 is used for detection. 75 ml of solution can be used for 3 blots.
- 4. The membrane is incubated for 30 min. in the antibody solution with gentle shaking.
  - 5. The membrane are washed twice in washing buffer with gentle shaking. About 250 mls is used per wash for 3 blots.
- 6. The blots are equilibrated for 2-5 min in 60 ml detection buffer.
  - 7. Dilute CSPD (1:200) in detection buffer. (This can be prepared ahead of time and stored in the dark at  $4^{\circ}$ C).
- The following steps must be done individually. Bags (one for detection and one for exposure) should be cut and ready before doing the following steps.

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- 8. The blot is carefully removed from the detection buffer and excess liquid removed without drying the membrane. The blot is immediately placed in a bag and 1.5 ml of CSPD solution is added. The CSPD solution can be spread over the membrane. Bubbles present at the edge and on the surface of the blot should be removed by gentle rubbing. The membrane is incubated for 5 min. in CSPD solution.
- 9. Excess liquid is removed and the membrane is blotted briefly (DNA side up) on Whatman 3MM paper. Do not let the membrane dry completely.
  - 10. Seal the damp membrane in a hybridization bag and incubate for 10 min at  $37\,^{\circ}\text{C}$  to enhance the luminescent reaction.
- 15 11. Expose for 2 hours at room temperature to X-ray film.

  Multiple exposures can be taken. Luminescence continues
  for at least 24 hours and signal intensity increases
  during the first hours.

## Example 2: Transformation of Carrot Cells

- 20 Transformation of plant cells can be accomplished by a number of methods, as described above. Similarly, a number of plant genera can be regenerated from tissue culture following transformation. Transformation and regeneration of carrot cells as described herein is illustrative.
- Single cell suspension cultures of carrot (Daucus carota) cells are established from hypocotyls of cultivar Early Nantes in  $B_5$  growth medium (O.L. Gamborg et al., Plant Physiol.  $\underline{45}$ :372 (1970)) plus 2,4-D and 15 mM CaCl<sub>2</sub> ( $B_5$  -44 medium) by methods known in the art. The suspension cultures are subcultured by adding 10 ml of the suspension culture to

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40 ml of  $B_5-44$  medium in 250 ml flasks every 7 days and are maintained in a shaker at 150 rpm at 27 °C in the dark.

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suspension culture cells are transformed with exogenous DNA as described by Z. Chen et al. Plant Mol. Bio. Briefly, 4-days post-subculture cells are 36:163 (1998). incubated with cell wall digestion solution containing 0.4 M sorbitol. 2% driselase, 5mM MES (2-[N-Morpholino] ethanesulfonic acid) pH 5.0 for 5 hours. The digested cells are pelleted gently at 60 xg for 5 min. and washed twice in W5 solution containing 154 mM NaCl, 5 mM KCl, 125 mM CaCl<sub>2</sub> and 5mM glucose, pH 6.0. The protoplasts are suspended in MC solution containing 5 mM MES, 20 mM CaCl<sub>2</sub>, 0.5 M mannitol, pH 5.7 and the protoplast density is adjusted to about 4 x  $10^6$ protoplasts per ml.

15-60  $\mu$ g of plasmid DNA is mixed with 0.9 ml The resulting suspension is mixed with 40% polyethylene glycol (MW 8000, PEG 8000), by gentle inversion a few times at room temperature for 5 to 25 min. Protoplast culture medium known in the art is added into the PEG-DNA-20 protoplast mixture. Protoplasts are incubated in the culture medium for 24 hour to 5 days and cell extracts can be used for assay of transient expression of the introduced gene. Alternatively, transformed cells can be used to produce transgenic callus, which in turn can be used to produce 25 transgenic plants, by methods known in the art. See, for example, Nomura and Komamine, Plt. Phys. 79:988-991 (1985), Identification and Isolation of Single Cells that Produce Somatic Embryos in Carrot Suspension Cultures.

The invention being thus described, it will be apparent 30 of ordinary skill in the art that modifications of the materials and methods for practicing the invention can be made. Such modifications are to be

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considered within the scope of the invention as defined by the following claims.

Each of the references from the patent and periodical literature cited herein is hereby expressly incorporated in its entirety by such citation.

110

```
REF TABLE 1
```

Maximum Length Sequence:

related to:

Clone IDs:

5

9581

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 1
  - Ceres seq id 1007546
- Alternative transcription start site(s) located in SEQ 10 ID NO 1:

-96, -51, 25, 27, 29, 55, 64

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 2
- Ceres seq id 1007547
  - Location of start within SEQ ID NO 1: at 1 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 2: at 32 aa.
- 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 1
    - gi No. 3879939
- 25 Description:
  - % Identity: 45.9
  - Alignment Length: 61
  - Location of Alignment in SEQ ID NO 2: from 48 to

108

30

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 3
  - Ceres seq id 1007548
  - Location of start within SEQ ID NO 1: at 100 nt.

35

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- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 2
    - gi No. 3879939
    - Description:
    - % Identity: 45.9
    - Alignment Length: 61
    - Location of Alignment in SEQ ID NO 3: from 15 to

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 4
  - Ceres seq id 1007549
- Location of start within SEQ ID NO 1: at 121 nt.

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```
(C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 3
 5
               - gi No. 3879939
               - Description:
               - % Identity: 45.9
               - Alignment Length: 61
     - Location of Alignment in SEQ ID NO 4: from 8 to 68
10
          (Ba) Polypeptide Activities: Similar to yeast membrane
    protein activities
15
    Maximum Length Sequence:
      related to:
     Clone IDs:
           9568
           402131
20
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEO ID NO 5
          - Ceres seq id 1007583
          - Alternative transcription start site(s) located in SEQ
     ID NO 5:
25
            2, 3, 4, 7, 10, 11, 12, 17, 42, 43, 300, 505
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 6
            - Ceres seq id 1007584
30
            - Location of start within SEQ ID NO 5: at 55 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - KH domain
35
               - Location within SEQ ID NO 6: from 47 to 95 aa.
            (Dp) Related Amino Acid Sequences
                - Alignment No. 4
                - gi No. 133940
40
                - Description:
               - % Identity: 75.1
                - Alignment Length: 250
                - Location of Alignment in SEQ ID NO 6: from 1 to
     249
45
         (B) Polypeptide Sequence
             - Pat. Appln. SEQ ID NO 7
             - Ceres seq id 1007585
             - Location of start within SEQ ID NO 5: at 184 nt.
```

```
(C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
               - KH domain
               - Location within SEQ ID NO 7: from 4 to 52 aa.
5
           (Dp) Related Amino Acid Sequences
               - Alignment No. 5
               - gi No. 133940
               - Description:
10
               - % Identity: 75.1
               - Alignment Length: 250
               - Location of Alignment in SEQ ID NO 7: from 1 to
     206
15
    Maximum Length Sequence:
      related to:
     Clone IDs:
           91769
     (Ac) cDNA Polynucleotide Sequence
20
          - Pat. Appln. SEQ ID NO 8
          - Ceres seq id 1008148
          - Alternative transcription start site(s) located in SEQ
     ID NO 8:
            -19,2,3,4,5,6,7,9,10,11,12,14
25
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 9
            - Ceres seq id 1008149
            - Location of start within SEQ ID NO 8: at 3 nt.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
               - Alignment No. 6
35
               - qi No. 4539292
               - Description:
               - % Identity:
                               89.9
               - Alignment Length: 181
               - Location of Alignment in SEQ ID NO 9: from 25 to
40
     203
         (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 10
            - Ceres seq id 1008150
45
            - Location of start within SEQ ID NO 8: at 75 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
50
                - Alignment No. 7
                - qi No. 4539292
```

113

```
- Description:
```

- % Identity: 89.9
- Alignment Length: 181
- Location of Alignment in SEQ ID NO 10: from 1 to

5 179

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 11
  - Ceres seq id 1008151
- 10 Location of start within SEQ ID NO 8: at 210 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
- Alignment No. 8
  - gi No. 4539292
  - Description:
  - % Identity: 89.9
  - Alignment Length: 181
- Location of Alignment in SEQ ID NO 11: from 1 to 134
- (Ba) Polypeptide Activities: Similar to 40S Ribosomal protein activities, and glycine rich RNA binding protein activities.

Maximum Length Sequence:

related to:

30 Clone IDs:

40

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 12
  - Ceres seq id 1008334
- Alternative transcription start site(s) located in SEQ ID NO 12:
  - -12,29,30
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 13
    - Ceres seq id 1008335
    - Location of start within SEQ ID NO 12: at 2 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 9
    - qi No. 3602948
    - Description:
- % Identity: 43.7
  - Alignment Length: 71

114

- Location of Alignment in SEQ ID NO 13: from 178 to 248  $\,$ 

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 14
  - Ceres seq id 1008336
  - Location of start within SEQ ID NO 12: at 59 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 14: at 22 aa.

(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

- (Dp) Related Amino Acid Sequences
  - Alignment No. 10
  - qi No. 3602948
  - Description:
  - % Identity: 43.7
  - Alignment Length: 71
  - Location of Alignment in SEQ ID NO 14: from 159
- 20 to 229

5

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- (Ba) Polypeptide Activities: Plant specific gene, Chloroplast specific gene.
- 25 Maximum Length Sequence:

related to:

Clone IDs:

8286

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 15
  - Ceres seq id 1008701
- Alternative transcription start site(s) located in SEQ ID NO 15:
  - -6, 2, 3, 5, 6, 7, 18, 24, 25, 28, 31, 33, 35, 37, 42, 50

(B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 16
- Ceres seq id 1008702
- Location of start within SEQ ID NO 15: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 11
    - qi No. 1083282
    - Description:
    - % Identity: 41.4
    - Alignment Length: 106
  - Location of Alignment in SEQ ID NO 16: from 23 to
- 50 121

```
115
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 17
            - Ceres seq id 1008703
            - Location of start within SEQ ID NO 15: at 67 nt.
 5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 12
10
               - qi No. 1083282
               - Description:
               - % Identity: 41.4
               - Alignment Length: 106
               - Location of Alignment in SEQ ID NO 17: from 1 to
15
     99
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 18
            - Ceres seq id 1008704
20
            - Location of start within SEQ ID NO 15: at 2 nt.
     (Ba) Polypeptide Activities: Similar to cytochrome C oxidase
     activities.
25
     Maximum Length Sequence:
      related to:
     Clone IDs:
           7792
30
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 19
          - Ceres seq id 1009003
          - Alternative transcription start site(s) located in SEQ
     ID NO 19:
35
            2,374
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 20
            - Ceres seq_id 1009004
40
            - Location of start within SEQ ID NO 19: at 48 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
45
                - Alignment No. 13
```

- gi No. 3582320
- Description:
- % Identity: 32.6
- Alignment Length: 44
- Location of Alignment in SEQ ID NO 20: from 51 to

WO 00/40695

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 21
            - Ceres seq id 1009005
 5
            - Location of start within SEQ ID NO 19: at 57 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
10
               - Alignment No. 14
               - gi No. 3582320
               - Description:
               - % Identity:
                               32.6
               - Alignment Length: 44
15
               - Location of Alignment in SEQ ID NO 21: from 48 to
     90
     (Ba) Polypeptide Activities: Similar to Ring-H2 Zinc Finger
     Protein activities.
20
     Maximum Length Sequence:
      related to:
     Clone IDs:
25
           7337
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 22
          - Ceres seq id 1009345
          - Alternative transcription start site(s) located in SEQ
30
     ID NO 22:
            2
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 23
35
            - Ceres seq id 1009346
            - Location of start within SEQ ID NO 22: at 50 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 23: at 22 aa.
40
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
                - Alignment No. 15
                - qi No. 3176705
45
                - Description:
                - % Identity:
                               53.3
                - Alignment Length: 75
                - Location of Alignment in SEQ ID NO 23: from 1 to
     75
50
         (B) Polypeptide Sequence
```

117

- Pat. Appln. SEQ ID NO 24
- Ceres seq id 1009347
- Location of start within SEQ ID NO 22: at 62 nt.
- Location of Signal Peptide Cleavage Site within SEQ
- 5 ID NO 24: at 18 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
- Alignment No. 16
  - gi No. 3176705
  - Description:
  - % Identity: 53.3
  - Alignment Length: 75
- Location of Alignment in SEQ ID NO 24: from 1 to 71
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot
- 20 specific gene, plant specific gene.

Maximum Length Sequence:

related to:

25 Clone IDs:

35

6349

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 25
  - Ceres seq id 1010140
- Alternative transcription start site(s) located in SEQ ID NO 25:
  - -31, -29, 4, 5, 6, 10, 17, 34, 41, 749
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 26
    - Ceres seq\_id 1010141
    - Location of start within SEQ ID NO 25: at 3 nt.
- (C) Nomination and Annotation of Domains within 40 Predicted Polypeptide(s)
  - Bacterial regulatory proteins, deoR family
  - Location within SEQ ID NO 26: from 57 to 95 aa.
  - (Dp) Related Amino Acid Sequences
- 45 Alignment No. 17
  - gi No. 3257798
  - Description:
  - % Identity: 40.6
  - Alignment Length: 234
- Location of Alignment in SEQ ID NO 26: from 56 to

118

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 27
            - Ceres seq id 1010142
 5
            - Location of start within SEQ ID NO 25: at 42 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Bacterial regulatory proteins, deoR family
10
               - Location within SEQ ID NO 27: from 44 to 82 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 18
               - qi No. 3257798
15
               - Description:
               - % Identity: 40.6
               - Alignment Length: 234
               - Location of Alignment in SEQ ID NO 27: from 43 to
     276
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 28
            - Ceres seq id 1010143
            - Location of start within SEQ ID NO 25: at 231 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 19
30
               - qi No. 3257798
               - Description:
               - % Identity: 40.6
               - Alignment Length: 234
               - Location of Alignment in SEQ ID NO 28: from 1 to
35
     213
     Maximum Length Sequence:
      related to:
     Clone IDs:
40
           6261
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 29
          - Ceres seq id 1010217
          - Alternative transcription start site(s) located in SEQ
45
     ID NO 29:
            2,5,15
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 30
50
            - Ceres seq id 1010218
```

- Location of start within SEQ ID NO 29: at 85 nt.

119 - Location of Signal Peptide Cleavage Site within SEQ ID NO 30: at 22 aa. (C) Nomination and Annotation of Domains within 5 Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 20 - qi No. 3341723 - Description: 10 - % Identity: 64.3 - Alignment Length: 131 - Location of Alignment in SEQ ID NO 30: from 1 to 118 15 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 31 - Ceres seq id 1010219 - Location of start within SEQ ID NO 29: at 118 nt. - Location of Signal Peptide Cleavage Site within SEQ 20 ID NO 31: at 13 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 25 - Alignment No. 21 - gi No. 3341723 - Description: - % Identity: 64.3 - Alignment Length: 131 30 - Location of Alignment in SEQ ID NO 31: from 1 to 107 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 32 - Ceres seq id 1010220 35 - Location of start within SEQ ID NO 29: at 121 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 40 - Alignment No. 22 - qi No. 3341723

- Description:
- % Identity: 64.3
- Alignment Length: 131
  - Location of Alignment in SEQ ID NO 32: from 1 to 106
- (Ba) Polypeptide Activities: Similar to Constans like 50 Protein activities and Zinc Finger Protein Activities.

```
Maximum Length Sequence:
      related to:
     Clone IDs:
 5
           6145
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 33
          - Ceres seq id 1010302
          - Alternative transcription start site(s) located in SEQ
     ID NO 33:
10
            -5, -3, -2, -1, 2, 3, 4, 5, 6, 7, 8, 10, 11, 12, 13, 15, 19, 23, 45, 349
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 34
15
            - Ceres seq id 1010303
            - Location of start within SEQ ID NO 33: at 59 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
20
               - Pathogenesis-related protein Bet v I family
               - Location within SEQ ID NO 34: from 5 to 155 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 23
25
               - qi No. 1321731
               - Description:
               - % Identity: 35.7
               - Alignment Length: 159
               - Location of Alignment in SEQ ID NO 34: from 5 to
30
     155
     Maximum Length Sequence:
      related to:
     Clone IDs:
35
           5180
     Public Genomic DNA:
           gi No: 4757410
           Predicted Exons:
              INTR 37202 ... 37397 OCKHAMG-CDNA
                     37493 ... 37825
40
              INTR
                                         OCKHAMG-CDNA
              TINI
                     37271 ... 37397
                                         OCKHAMG-CDS
              TERM
                     37493 ... 37704
                                         OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
45
          - Pat. Appln. SEQ ID NO 35
          - Ceres seq id 1010815
          - Alternative transcription start site(s) located in SEQ
     ID NO 35:
            15, 16, 17, 18, 19, 29, 31, 34
50
        (B) Polypeptide Sequence
```

121

- Pat. Appln. SEQ ID NO 36
- Ceres seq id 1010816
- Location of start within SEQ ID NO 35: at 70 nt.
- 5 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 24
    - gi No. 2879811
- 10 Description:
  - % Identity: 88.4
  - Alignment Length: 112
  - Location of Alignment in SEQ ID NO 36: from 1 to

112

15

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 37
  - Ceres seq id 1010817
  - Location of start within SEQ ID NO 35: at 133 nt.

20

25

- - (Dp) Related Amino Acid Sequences
    - Alignment No. 25
    - gi No. 2879811
      - Description:
      - % Identity: 88.4
      - Alignment Length: 112
      - Location of Alignment in SEQ ID NO 37: from 1 to
- 30 91
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 38
  - Ceres seq id 1010818
- 35 Location of start within SEQ ID NO 35: at 257 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 38: at 26 aa.
- (Ba) Polypeptide Activities: Similar to ribosomal protein actitivies.

Maximum Length Sequence:

related to:

Clone IDs:

45 42842

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 39
  - Ceres seq id 1011437
- Alternative transcription start site(s) located in SEQ
- 50 ID NO 39:

-28

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 40
            - Ceres seq id 1011438
 5
            - Location of start within SEQ ID NO 39: at 2 nt.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 41
            - Ceres seq id 1011439
10
            - Location of start within SEQ ID NO 39: at 1 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 41: at 29 aa.
           (C) Nomination and Annotation of Domains within
15
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 26
               - qi No. 3334271
               - Description:
               - % Identity: 29.6
20
               - Alignment Length: 54
               - Location of Alignment in SEQ ID NO 41: from 13 to
     65
25
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 42
            - Ceres seq id 1011440
            - Location of start within SEQ ID NO 39: at 28 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 42: at 20 aa.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
35
               - Alignment No. 27
               - qi No. 3334271
               - Description:
               - % Identity:
                              29.6
               - Alignment Length: 54
40
               - Location of Alignment in SEQ ID NO 42: from 4 to
     56
     (Ba) Polypeptide Activities: Similar to NADH Oxidoxductase
     MWFE Subunit Protein Activities.
45
     Maximum Length Sequence:
      related to:
     Clone IDs:
50
           42475
     (Ac) cDNA Polynucleotide Sequence
```

123

- Pat. Appln. SEQ ID NO 43
- Ceres seq id 1011616
- Alternative transcription start site(s) located in SEQ ID NO 43:

5 3,5,476

25

35

50

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 44
  - Ceres seq id 1011617
- Location of start within SEQ ID NO 43: at 115 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
- Alignment No. 28
  - qi No. 3859560
  - Description:
  - % Identity: 36.2
  - Alignment Length: 225
- Location of Alignment in SEQ ID NO 44: from 31 to 248
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 45
  - Ceres seq\_id 1011618
    - Location of start within SEQ ID NO 43: at 406 nt.
    - (C) Nomination and Annotation of Doma, s within Predicted Polypeptide(s)
- 30 (Dp) Related Amino Acid Sequences
  - Alignment No. 29
  - qi No. 3859560
  - Description:
  - % Identity: 36.2
  - Alignment Length: 225
    - Location of Alignment in SEQ ID NO 45: from 1 to 151
- (Ba) Polypeptide Activities: Similar to acycl-protein 40 thioosterases protein activities, calcium independent phospholipase A2 activities, and carboxylesterase activities.
- 45 Maximum Length Sequence:

related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 46
    - Ceres seq id 1011631

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 47
            - Ceres seq id 1011632
 5
            - Location of start within SEQ ID NO 46: at 3 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
10
               - Alignment No. 30
               - qi No. 3618318
               - Description:
                              72.7
               - % Identity:
15
               - Alignment Length: 44
               - Location of Alignment in SEQ ID NO 47: from 91 to
     134
        (B) Polypeptide Sequence
20
            - Pat. Appln. SEQ ID NO 48
            - Ceres seq id 1011633
            - Location of start within SEQ ID NO 46: at 9 nt.
           (C) Nomination and Annotation of Domains within
25
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 31
               - qi No. 3618318
               - Description:
               - % Identity: 72.7
30
               - Alignment Length: 44
               - Location of Alignment in SEQ ID NO 48: from 89 to
     132
35
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 49
            - Ceres seq id 1011634
            - Location of start within SEQ ID NO 46: at 15 nt.
40
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 32
               - qi No. 3618318
               - Description:
45
               - % Identity: 72.7
               - Alignment Length: 44
               - Location of Alignment in SEQ ID NO 49: from 87 to
     130
50
```

125

(Ba) Polypeptide Activities: Similar to Constans protein activities, and zinc finger protein activities.

Maximum Length Sequence:

related to:

Clone IDs:

5

20

25

42240

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 50
- 10 Ceres seq id 1011714
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 51
- Ceres seq id 1011715
  - Location of start within SEQ ID NO 50: at 2 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Bacterial mutT protein
      - Location within SEQ ID NO 51: from 26 to 67 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 33
      - qi No. 2129134
      - Description:
      - % Identity: 40.8
      - Alignment Length: 121
      - Location of Alignment in SEQ ID NO 51: from 12 to
- 30 131
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 52
    - Ceres seq id 1011716
- Location of start within SEQ ID NO 50: at 14 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Bacterial mutT protein
- Location within SEQ ID NO 52: from 22 to 63 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 34
    - gi No. 2129134
- 45 Description:
  - % Identity: 40.8
  - Alignment Length: 121
  - Location of Alignment in SEQ ID NO 52: from 8 to

127 50

(B) Polypeptide Sequence

126

- Pat. Appln. SEQ ID NO 53
- Ceres seq id 1011717
- Location of start within SEQ ID NO 50: at 185 nt.
- 5 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 35
    - qi No. 2129134
- 10 Description:
  - % Identity: 40.8
  - Alignment Length: 121
  - Location of Alignment in SEQ ID NO 53: from 1 to

70

15

30

40

Maximum Length Sequence:

related to:

Clone IDs:

- 20 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 54
  - Ceres seq id 1011784
  - Alternative transcription start site(s) located in SEQ ID NO 54:
- 25 -15, -4, 7, 402
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 55
    - Ceres seq id 1011785
  - Location of start within SEQ ID NO 54: at 1 nt.
    - Location of Signal Peptide Cleavage Site within SEQ ID NO  $55\colon$  at 42 aa.
- (C) Nomination and Annotation of Domains within
- 35 Predicted Polypeptide(s)
  - Plant lipid transfer protein family
  - Location within SEQ ID NO 55: from 45 to 108 aa.
  - (Dp) Related Amino Acid Sequences
  - Alignment No. 36
    - qi No. 543565
    - Description:
    - % Identity: 56.5
    - Alignment Length: 85
- Location of Alignment in SEQ ID NO 55: from 29 to
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 56
- 50 Ceres seq id 1011786
  - Location of start within SEQ ID NO 54: at 49 nt.

127 - Location of Signal Peptide Cleavage Site within SEQ ID NO 56: at 26 aa. (C) Nomination and Annotation of Domains within 5 Predicted Polypeptide(s) - Plant lipid transfer protein family - Location within SEQ ID NO 56: from 29 to 92 aa. (Dp) Related Amino Acid Sequences - Alignment No. 37 10 - qi No. 543565 - Description: - % Identity: 56.5 - Alignment Length: 85 15 - Location of Alignment in SEQ ID NO 56: from 13 to 94 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 57 20 - Ceres seq id 1011787 - Location of start within SEQ ID NO 54: at 3 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) 25 (Dp) Related Amino Acid Sequences Maximum Length Sequence: related to: Clone IDs: 30 41992 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 58 - Ceres seq id 1011820 - Alternative transcription start site(s) located in SEQ 35 ID NO 58: -40,37(B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 59 40 - Ceres seq id 1011821

- Location of start within SEQ ID NO 58: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 45 (Dp) Related Amino Acid Sequences
  - Alignment No. 38
  - qi No. 3417418
  - Description:
  - % Identity: 23.6
- Alignment Length: 207

128

- Location of Alignment in SEQ ID NO 59: from 24 to 226

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 60
  - Ceres seq id 1011822
  - Location of start within SEQ ID NO 58: at 13 nt.
- (C) Nomination and Annotation of Domains within 10 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 39
    - qi No. 3417418
    - Description:
- % Identity: 23.6
  - Alignment Length: 207
  - Location of Alignment in SEQ ID NO 60: from 20 to
- 20 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 61
  - Ceres seq id 1011823
  - Location of start within SEQ ID NO 58: at 151 nt.
- 25 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 40
    - gi No. 3417418
  - Description:
    - % Identity: 23.6
    - Alignment Length: 207
    - Location of Alignment in SEQ ID NO 61: from 1 to

176

30

35

40

222

5

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

Maximum Length Sequence:

related to:

Clone IDs:

- 45 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 62
  - Ceres seq id 1011874
- 50 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 63

129

- Ceres seq id 1011875
- Location of start within SEQ ID NO 62: at 1 nt.
- (C) Nomination and Annotation of Domains within
  5 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 41
    - qi No. 2911044
    - Description:
- % Identity: 78.7
  - Alignment Length: 95
  - Location of Alignment in SEQ ID NO 63: from 28 to 121
- (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

Maximum Length Sequence:

related to:

Clone IDs:

20 41682

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 64
  - Ceres seq id 1011981

25

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 65
  - Ceres seq id 1011982
  - Location of start within SEQ ID NO 64: at 68 nt.

30

35

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 42
    - gi No. 4115355
    - Description:
    - % Identity: 100
    - Alignment Length: 52
    - Location of Alignment in SEQ ID NO 65: from 1 to

40 52

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 66
  - Ceres seq id 1011983
- 45 Location of start within SEQ ID NO 64: at 3 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous} % \begin{subarray}{ll} \end{subarray} % \begin{s$ 
    - (Dp) Related Amino Acid Sequences

50

(B) Polypeptide Sequence

130

- Pat. Appln. SEQ ID NO 67

- Ceres seq id 1011984

- Location of start within SEQ ID NO 64: at 483 nt.

- Location of Signal Peptide Cleavage Site within SEQ

5 ID NO 67: at 19 aa.

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

10

Maximum Length Sequence:

related to:

Clone IDs:

15 38470

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 68
  - Ceres seq id 1014547
- Alternative transcription start site(s) located in SEQ 20 ID NO 68:

-39, -2, -1, 2, 3, 7, 8, 9, 10

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 69
- 25 Ceres seq id 1014548
  - Location of start within SEQ ID NO 68: at 1 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 69: at 32 aa.
- 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Plant lipid transfer protein family
  - Location within SEQ ID NO 69: from 44 to 110 aa.
- 35 (Dp) Related Amino Acid Sequences
  - Alignment No. 43
  - gi No. 3062791
  - Description:
  - % Identity: 72.2
- Alignment Length: 90
  - Location of Alignment in SEQ ID NO 69: from 21 to
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 70
    - Ceres seq id 1014549
    - Location of start within SEQ ID NO 68: at 25 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 70: at 24 aa.

50

45

131

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Plant lipid transfer protein family
               - Location within SEQ ID NO 70: from 36 to 102 aa.
 5
           (Dp) Related Amino Acid Sequences
               - Alignment No. 44
               - qi No. 3062791
               - Description:
               - % Identity:
10
                              72.2
               - Alignment Length: 90
               - Location of Alignment in SEQ ID NO 70: from 13 to
     102
     Maximum Length Sequence:
15
      related to:
     Clone IDs:
           38004
     (Ac) cDNA Polynucleotide Sequence
20
          - Pat. Appln. SEQ ID NO 71
          - Ceres seq id 1014995
          - Alternative transcription start site(s) located in SEQ
     ID NO 71:
            2,3,4
25
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 72
            - Ceres seq id 1014996
            - Location of start within SEQ ID NO 71: at 2 nt.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Aminotransferase class IV
               - Location within SEQ ID NO 72: from 71 to 334 aa.
35
            (Dp) Related Amino Acid Sequences
               - Alignment No. 45
               - gi No. 3540183
               - Description:
40
               - % Identity: 54.7
               - Alignment Length: 287
               - Location of Alignment in SEQ ID NO 72: from 56 to
     341
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 73
            - Ceres seq id 1014997
            - Location of start within SEQ ID NO 71: at 65 nt.
```

(C) Nomination and Annotation of Domains within

50

Predicted Polypeptide(s)

132

```
- Aminotransferase class IV
```

- Location within SEQ ID NO 73: from 50 to 313 aa.

(Dp) Related Amino Acid Sequences

- Alignment No. 46

- gi No. 3540183
- Description:
- % Identity: 54.7
- Alignment Length: 287
- Location of Alignment in SEQ ID NO 73: from 35 to 320
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 74
- Ceres seq\_id 1014998
  - Location of start within SEQ ID NO 71: at 167 nt.
  - (C) Nomination and Annotation of Domains within  $Predicted\ Polypeptide(s)$
  - Aminotransferase class IV
    - Location within SEQ ID NO 74: from 16 to 279 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 47
      - qi No. 3540183
      - Description:
      - % Identity: 54.7
      - Alignment Length: 287
      - Location of Alignment in SEQ ID NO 74: from 1 to
- 30 286

5

20

25

Maximum Length Sequence:

related to:

Clone IDs:

35 37701

Public Genomic DNA:

gi No: 4699904

Predicted Exons:

SINGLE 38530 ... 37988 OCKHAMG-CDS

- 40 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 75
  - Ceres seq id 1015323
  - Alternative transcription start site(s) located in SEQ ID NO 75:
- 45 -1,5,6,12
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 76
    - Ceres seq id 1015324
- 50 Location of start within SEQ ID NO 75: at 59 nt.

133

- Location of Signal Peptide Cleavage Site within SEQ ID NO 76: at 25 aa.

- (C) Nomination and Annotation of Domains within
  Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 48
    - qi No. 3860308
    - Description:
- % Identity: 44.5
  - Alignment Length: 140
  - Location of Alignment in SEQ ID NO 76: from 56 to
- 15 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

20 Maximum Length Sequence:

related to:

Clone IDs:

364

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 77
  - Ceres seq id 1016486
- Alternative transcription start site(s) located in SEQ ID NO 77:

17, 19, 20, 21, 22, 23, 29, 35, 38

30

40

45

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 78
  - Ceres seq id 1016487
  - Location of start within SEQ ID NO 77: at 1 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 78: at 46 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Gamma-thionins family
      - Location within SEQ ID NO 78: from 55 to 101 aa.
      - (Dp) Related Amino Acid Sequences
        - Alignment No. 49
        - qi No. 4038039
        - Description:
        - % Identity: 100
        - Alignment Length: 77
        - Location of Alignment in SEQ ID NO 78: from 25 to
- 50 101

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 79
            - Ceres seq id 1016488
            - Location of start within SEQ ID NO 77: at 73 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
5
     ID NO 79: at 22 aa.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Gamma-thionins family
10
               - Location within SEQ ID NO 79: from 31 to 77 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 50
15
               - qi No. 4038039
               - Description:
               - % Identity:
                              100
               - Alignment Length: 77
               - Location of Alignment in SEQ ID NO 79: from 1 to
     77
2.0
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 80
            - Ceres seq id 1016489
25
            - Location of start within SEQ ID NO 77: at 118 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Gamma-thionins family
30
               - Location within SEQ ID NO 80: from 16 to 62 aa.
            (Dp) Related Amino Acid Sequences
               - Alignment No. 51
               - gi No. 4038039
35
               - Description:
               - % Identity:
                               100
               - Alignment Length: 77
                - Location of Alignment in SEQ ID NO 80: from 1 to
     62
40
     Maximum Length Sequence:
      related to:
     Clone IDs:
           33891
45
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 81
          - Ceres seq id 1018341
          - Alternative transcription start site(s) located in SEQ
     ID NO 81:
50
             4
```

```
135
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 82
            - Ceres seq id 1018342
            - Location of start within SEQ ID NO 81: at 71 nt.
 5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 52
10
               - qi No. 1064887
               - Description:
               - % Identity:
               - Alignment Length: 65
               - Location of Alignment in SEQ ID NO 82: from 2 to
15
     66
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 83
            - Ceres seq id 1018343
20
            - Location of start within SEQ ID NO 81: at 143 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
25
               - Alignment No. 53
               - qi No. 1064887
               - Description:
               - % Identity:
                               60
               - Alignment Length: 65
30
               - Location of Alignment in SEQ ID NO 83: from 1 to
     42
         (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 84
35
            - Ceres seq id 1018344
            - Location of start within SEQ ID NO 81: at 146 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
40
            (Dp) Related Amino Acid Sequences
                - Alignment No. 54
                - qi No. 1064887
                - Description:
                - % Identity:
                               60
                - Alignment Length: 65
45
                - Location of Alignment in SEQ ID NO 84: from 1 to
```

(Ba) Polypeptide Activities: Similar to pollen coat protein activities and LEA protein activities.

41

```
136
    Maximum Length Sequence:
      related to:
    Clone IDs:
           33828
     (Ac) cDNA Polynucleotide Sequence
5
          - Pat. Appln. SEQ ID NO 85
          - Ceres seg id 1018382
          - Alternative transcription start site(s) located in SEQ
     ID NO 85:
10
            2,4,5,6,7,8,9,10,11,14
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 86
            - Ceres seq id 1018383
15
            - Location of start within SEQ ID NO 85: at 22 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 86: at 24 aa.
           (C) Nomination and Annotation of Domains within
20
     Predicted Polypeptide(s)
               - Plant lipid transfer protein family
               - Location within SEQ ID NO 86: from 28 to 115 aa.
           (Dp) Related Amino Acid Sequences
25
               - Alignment No. 55
               - qi No. 899224
               - Description:
               - % Identity: 78.2
               - Alignment Length: 119
30
               - Location of Alignment in SEQ ID NO 86: from 1 to
     119
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 87
35
            - Ceres seq id 1018384
            - Location of start within SEQ ID NO 85: at 73 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
40
               - Plant lipid transfer protein family
               - Location within SEQ ID NO 87: from 11 to 98 aa.
            (Dp) Related Amino Acid Sequences
                - Alignment No. 56
45
               - gi No. 899224
               - Description:
                - % Identity: 78.2
                - Alignment Length: 119
```

- Location of Alignment in SEQ ID NO 87: from 1 to

50

137

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 88
            - Ceres seq id 1018385
            - Location of start within SEQ ID NO 85: at 3 nt.
 5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
10
     Maximum Length Sequence:
      related to:
     Clone IDs:
           30349
     (Ac) cDNA Polynucleotide Sequence
15
          - Pat. Appln. SEQ ID NO 89
          - Ceres seq id 1020666
          - Alternative transcription start site(s) located in SEQ
     ID NO 89:
            33, 35, 39, 40, 42, 43, 44, 45, 64, 173
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 90
            - Ceres seq id 1020667
            - Location of start within SEQ ID NO 89: at 118 nt.
25
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein S7e
               - Location within SEQ ID NO 90: from 7 to 187 aa.
30
            (Dp) Related Amino Acid Sequences
                - Alignment No. 57
               - qi No. 3851636
               - Description:
35
               - % Identity:
                               77.4
               - Alignment Length: 190
               - Location of Alignment in SEQ ID NO 90: from 1 to
     190
40
         (B) Polypeptide Sequence
             - Pat. Appln. SEQ ID NO 91
             - Ceres seq id 1020668
             - Location of start within SEQ ID NO 89: at 271 nt.
45
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
                - Ribosomal protein S7e
                - Location within SEQ ID NO 91: from 1 to 136 aa.
50
            (Dp) Related Amino Acid Sequences
```

- Alignment No. 58

138

```
- gi No. 3851636
```

- Description:
- % Identity: 77.4
- Alignment Length: 190
- 5 Location of Alignment in SEQ ID NO 91: from 1 to

139

Maximum Length Sequence:

related to:

10 Clone IDs:

30113

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 92
  - Ceres seq id 1020784

15

30

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 93
  - Ceres seq id 1020785
- Location of start within SEQ ID NO 92: at 60 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 93: at 25 aa.
- (C) Nomination and Annotation of Domains within
- 25 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 59
    - qi No. 3860308
    - Description:
    - % Identity: 44.5
    - Alignment Length: 140
    - Location of Alignment in SEQ ID NO 93: from 56 to

175

35
(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

40 Maximum Length Sequence:

related to:

Clone IDs:

29120

Public Genomic DNA:

45 gi No: 5822667

Predicted Exons:

INTR 68772 ... 69532 OCKHAMG-CDNA

SINGLE 68846 ... 69325 OCKHAMG-CDS

gi No: 6041831 Predicted Exons:

139

INTR 63702 ... 64462 OCKHAMG-CDNA

SINGLE 63776 ... 64255 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 94
- Ceres seq id 1021525
- Alternative transcription start site(s) located in SEQ ID NO 94:

25, 26, 27, 28, 29, 35, 36, 39, 51, 53, 54, 68

10

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 95
  - Ceres seq id 1021526
  - Location of start within SEQ ID NO 94: at 75 nt.

15

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 60

20 - qi No. 4388980

- Description:
- % Identity: 29.1
- Alignment Length: 110
- Location of Alignment in SEQ ID NO 95: from 47 to

25 156

(Ba) Polypeptide Activities: Similar to adrenodoxi precursor protein activities and adrenal ferredoxin activities.

30

Maximum Length Sequence:

related to:

Clone IDs:

35 2891

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 96
  - Ceres seq id 1021563
- Alternative transcription start site(s) located in SEQ 40 ID NO 96:

16,28,29,30,31,35,36,43,74,77,80,88,89,90,95

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 97
- 45 Ceres seq id 1021564
  - Location of start within SEQ ID NO 96: at 2 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 50 (Dp) Related Amino Acid Sequences
  - Alignment No. 61

140

- qi No. 2493089
- Description:
- % Identity: 29.7
- Alignment Length: 77
- Location of Alignment in SEQ ID NO 97: from 52 to 125
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 98
- Ceres seq\_id 1021565
  - Location of start within SEQ ID NO 96: at 116 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 62
    - qi No. 2493089
    - Description:
    - % Identity: 29.7
  - Alignment Length: 77
    - Location of Alignment in SEQ ID NO 98: from 14 to 87
- (Ba) Polypeptide Activities: Similar to ATPK-mouse 25 activities, ATP synthase activities, and mitochondrial F-Chain activities.

Maximum Length Sequence:

related to:

30 Clone IDs:

15

20

45

28979

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 99
  - Ceres seq id 1021576
- Alternative transcription start site(s) located in SEQ ID NO 99:

-13,-

11,3,5,7,11,12,13,15,24,26,28,29,31,34,35,36,37,38,39

40 43,44,45,46,47,48,50,51,52,56,58,59,61,67,68,70,75,82,83,91,1

244,313,318

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 100
  - Ceres seq id 1021577
  - Location of start within SEQ ID NO 99: at 109 nt.
- (C) Nomination and Annotation of Domains within 50 Predicted Polypeptide(s)
  - Photosystem I psaG / psaK

141

- Location within SEQ ID NO 100: from 50 to 130 aa.

- (Dp) Related Amino Acid Sequences
  - Alignment No. 63
  - qi No. 3885511
  - Description:
  - % Identity: 81.1
  - Alignment Length: 128
  - Location of Alignment in SEQ ID NO 100: from 1 to

127

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 101
  - Ceres seq id 1021578
  - Location of start within SEQ ID NO 99: at 121 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Photosystem I psaG / psaK
  - Location within SEQ ID NO 101: from 46 to 126 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 64
    - qi No. 3885511
    - Description:
    - % Identity: 81.1
    - Alignment Length: 128
    - Location of Alignment in SEQ ID NO 101: from 1 to

123

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 102
  - Ceres seq id 1021579
  - Location of start within SEQ ID NO 99: at 124 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Photosystem I psaG / psaK
  - Location within SEQ ID NO 102: from 45 to 125 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 65
    - qi No. 3885511
    - Description:
    - % Identity: 81.1
    - Alignment Length: 128
    - Location of Alignment in SEQ ID NO 102: from 1 to

122

0 Maximum Length Sequence:
 related to:

)

142

Clone IDs:

28177

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 103
  - Ceres seq id 1021927
- Alternative transcription start site(s) located in SEQ ID NO 103:

2, 12, 20, 23, 29, 36, 46, 47, 53

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 104
  - Ceres seq id 1021928
  - Location of start within SEQ ID NO 103: at 67 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 66
    - qi No. 4263779
    - Description:
    - % Identity: 28.2
    - Alignment Length: 177
    - Location of Alignment in SEQ ID NO 104: from 14

to 182

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 105
  - Ceres seq\_id 1021929
  - Location of start within SEQ ID NO 103: at 172 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 105: at 23 aa.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\cdot$ 
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 67
    - qi No. 4263779
    - Description:
    - % Identity: 28.2
    - Alignment Length: 177
    - Location of Alignment in SEQ ID NO 105: from 1 to

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 106
  - Ceres seq id 1021930
  - Location of start within SEQ ID NO 103: at 220 nt.
- (C) Nomination and Annotation of Domains within
  O Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences

```
143
               - Alignment No. 68
               - gi No. 4263779
               - Description:
               - % Identity:
                              28.2
5
               - Alignment Length: 177
               - Location of Alignment in SEQ ID NO 106: from 1 to
     131
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
10
     Maximum Length Sequence:
      related to:
     Clone IDs:
15
           2807
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 107
          - Ceres seq id 1021945
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 108
            - Ceres seq id 1021946
            - Location of start within SEQ ID NO 107: at 1 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 69
30
               - gi No. 3334132
               - Description:
               - % Identity:
                              27.4
               - Alignment Length: 114
               - Location of Alignment in SEQ ID NO 108: from 28
35
     to 136
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 109
            - Ceres seq id 1021947
40
            - Location of start within SEQ ID NO 107: at 163 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
               - Alignment No. 70
45
               - gi No. 3334132
               - Description:
               - % Identity:
                              27.4
```

- Alignment Length: 114

- Location of Alignment in SEQ ID NO 109: from 1 to

82

144

(Ba) Polypeptide Activities: Similar to C21 ORF4 - membrane protein activities 5 Maximum Length Sequence: related to: Clone IDs: 27792 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 110 10 - Ceres seq id 1022170 - Alternative transcription start site(s) located in SEQ ID NO 110: -4, -1, 32, 6815 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 111 - Ceres seq id 1022171 - Location of start within SEQ ID NO 110: at 92 nt. 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 71 25 - gi No. 1173456 - Description: - % Identity: 54.7 - Alignment Length: 129 - Location of Alignment in SEQ ID NO 111: from 4 to 30 131 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 112 - Ceres seq id 1022172 35 - Location of start within SEQ ID NO 110: at 191 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 40 - Alignment No. 72 - qi No. 1173456 - Description: - % Identity: 54.7 - Alignment Length: 129 - Location of Alignment in SEQ ID NO 112: from 1 to 45 98 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 113

- Location of start within SEQ ID NO 110: at 1 nt.

- Ceres seq id 1022173

145

- Location of Signal Peptide Cleavage Site within SEQ ID NO 113: at 19  $\ensuremath{\mathrm{aa}}$  .

(Ba) Polypeptide Activities: Similar to small nuclear ribonucleoprotein activities.

Maximum Length Sequence:

related to:

Clone IDs:

10 27167

Public Genomic DNA:

gi No: 3046850

Predicted Exons:

INIT 45217 ... 45131 OCKHAMG-CDS INTR 44695 ... 44629 OCKHAMG-CDS TERM 44554 ... 44286 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 114
  - Ceres seq id 1022554
- Alternative transcription start site(s) located in SEQ ID NO 114:

-49,-

19,7,22,23,28,29,30,31,32,33,35,36,37,51,57,67,75,81,82

25

15

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 115
  - Ceres seq id 1022555
  - Location of start within SEQ ID NO 114: at 107 nt.

30

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- Heme-binding domain in cytochrome b5 and oxidoreductases
- Location within SEQ ID NO 115: from 7 to 84 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 73
    - gi No. 4240122
  - Description:
    - % Identity: 100
    - Alignment Length: 140
  - Location of Alignment in SEQ ID NO 115: from 1 to 140

45

40

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 116
  - Ceres seq\_id 1022556
  - Location of start within SEQ ID NO 114: at 317 nt.

146

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 74
 5
               - qi No. 4240122
               - Description:
               - % Identity: 100
               - Alignment Length: 140
               - Location of Alignment in SEQ ID NO 116: from 1 to
10
    70
     Maximum Length Sequence:
      related to:
     Clone IDs:
           27109
15
     Public Genomic DNA:
           qi No: 6449507
           Predicted Exons:
              INIT
                     94711 ... 94519 OCKHAMG-CDS
                     94417 ... 94326 OCKHAMG-CDS
20
              INTR
                     94249 ... 94131 OCKHAMG-CDS
              INTR
              TERM 94046 ... 93968
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 117
25
          - Ceres seq id 1022594
          - Alternative transcription start site(s) located in SEQ
     ID NO 117:
            2, 15, 24, 25, 66, 69, 72, 74
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 118
            - Ceres seq id 1022595
            - Location of start within SEQ ID NO 117: at 1 nt.
           (C) Nomination and Annotation of Domains within
35
     Predicted Polypeptide(s)
               - Universal stress protein family
               - Location within SEQ ID NO 118: from 98 to 183 aa.
40
           (Dp) Related Amino Acid Sequences
               - Alignment No. 75
               - qi No. 2160182
               - Description:
               - % Identity:
                              37.5
45
               - Alignment Length: 160
               - Location of Alignment in SEQ ID NO 118: from 39
     to 186
         (B) Polypeptide Sequence
50
            - Pat. Appln. SEQ ID NO 119
```

- Ceres seq id 1022596

147

- Location of start within SEQ ID NO 117: at 82 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Universal stress protein family
    - Location within SEQ ID NO 119: from 71 to 156 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 76
    - qi No. 2160182
    - Description:
    - % Identity: 37.5
    - Alignment Length: 160
    - Location of Alignment in SEQ ID NO 119: from 12
- 15 to 159

5

10

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 120
  - Ceres seq id 1022597
- 20 Location of start within SEQ ID NO 117: at 106 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Universal stress protein family
- Location within SEQ ID NO 120: from 63 to 148 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 77
    - gi No. 2160182
- 30 Description:
  - % Identity: 37.5
  - Alignment Length: 160
  - Location of Alignment in SEQ ID NO 120: from 4 to

151

35

(Ba) Polypeptide Activities: Similar to protein in methanobacterium thermoautotrophicum activities.

Maximum Length Sequence:

40 related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 121
- 45 Ceres seq id 1022621
  - Alternative transcription start site(s) located in SEQ ID NO 121:
    - 2,7,9,13,35,38,45,57
- 50 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 122

148

- Ceres seq id 1022622
- Location of start within SEQ ID NO 121: at 86 nt.
- (C) Nomination and Annotation of Domains within
  5 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 78
    - qi No. 3256599
    - Description:
    - % Identity: 32
    - Alignment Length: 128
  - Location of Alignment in SEQ ID NO 122: from 7 to 128
- 15 (Ba) Polypeptide Activities: Similar to structural cell wall protein activities, and larval gene protein in the Fruit fly activities.
- 20 Maximum Length Sequence:

related to:

Clone IDs:

23518

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 123
  - Ceres seq id 1024375
- Alternative transcription start site(s) located in SEQ ID NO 123:
  - 7,8,10,17,18,24,35,41,42,43,44,46,48,52,54,59

30

25

10

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 124
  - Ceres seq id 1024376
  - Location of start within SEQ ID NO 123: at 130 nt.

35

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S7e
  - Location within SEQ ID NO 124: from 7 to 187 aa.

40

45

- (Dp) Related Amino Acid Sequences
  - Alignment No. 79
  - qi No. 3851636
  - Description:
  - % Identity: 76.6
  - Alignment Length: 188
  - Location of Alignment in SEQ ID NO 124: from 1 to

188

50 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 125

149

- Ceres seq id 1024377
- Location of start within SEQ ID NO 123: at 283 nt.
- (C) Nomination and Annotation of Domains within
  5 Predicted Polypeptide(s)
  - Ribosomal protein S7e
  - Location within SEQ ID NO 125: from 1 to 136 aa.
  - (Dp) Related Amino Acid Sequences
- Alignment No. 80
  - qi No. 3851636
  - Description:
  - % Identity: 76.6
  - Alignment Length: 188
- Location of Alignment in SEQ ID NO 125: from 1 to

Maximum Length Sequence:

related to:

20 Clone IDs:

30

40

92

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 126
  - Ceres seq id 1024535
- Alternative transcription start site(s) located in SEQ ID NO 126:
  - 2,8,11,31,46,47,48
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 127
    - Ceres seq id 1024536
    - Location of start within SEQ ID NO 126: at 115 nt.
- (C) Nomination and Annotation of Domains within
- 35 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 81
    - gi No. 2621731
    - Description:
    - % Identity: 35.2
    - Alignment Length: 88
  - Location of Alignment in SEQ ID NO 127: from 5 to
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 128
  - Ceres seq id 1024537
  - Location of start within SEQ ID NO 126: at 3 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

150

(Dp) Related Amino Acid Sequences

```
(B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 129
- Ceres seq id 1024538
- Location of start within SEQ ID NO 126: at 253 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 10 (Dp) Related Amino Acid Sequences
  - Alignment No. 82
  - qi No. 2621731
  - Description:
  - % Identity: 35.2
- Alignment Length: 88
  - Location of Alignment in SEQ ID NO 129: from 1 to 46

26250

(Ba) Polypeptide Activities: Similar to small nuclear ribonucleoprotein activities.

Maximum Length Sequence:

related to:

25 Clone IDs:

5

21228

Public Genomic DNA:

gi No: 4539402

Predicted Exons: 30 INIT 36726

30	T $N$ $T$ $T$ .	36/26	 36352	GENBANK
	INTR	36257	 36135	GENBANK
	TERM	35555	 35469	GENBANK

INTR 36791 ... 36352 OCKHAMG-CDNA
INTR 36257 ... 36135 OCKHAMG-CDNA
INTR 35555 ... 35325 OCKHAMG-CDNA

INIT 36726 ... 36352 OCKHAMG-CDS INTR 36257 ... 36135 OCKHAMG-CDS 40 TERM 35555 ... 35469 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 130
- Ceres seq id 1025683
- Alternative transcription start site(s) located in SEQ 45 ID NO 130:

2,8,22

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 131
- 50 Ceres seq id 1025684
  - Location of start within SEQ ID NO 130; at 3 nt.

	<ul><li>(C) Nomination and Annotation of Domains within</li><li>Predicted Polypeptide(s)</li><li>- Ribosomal protein L6</li></ul>
5	- Location within SEQ ID NO 131: from 34 to 216 aa.
10	(Dp) Related Amino Acid Sequences - Alignment No. 83 - gi No. 266945 - Description:
	- % Identity: 84 - Alignment Length: 194 - Location of Alignment in SEQ ID NO 131: from 23 to 216
15	(B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 132 - Ceres seq_id 1025685
20	- Location of start within SEQ ID NO 130: at 69 nt.  (C) Nomination and Annotation of Domains within
	Predicted Polypeptide(s)  - Ribosomal protein L6  - Location within SEQ ID NO 132: from 12 to 194 aa.
25	(Dp) Related Amino Acid Sequences - Alignment No. 84 - gi No. 266945 - Description:
30	- % Identity: 84 - Alignment Length: 194 - Location of Alignment in SEQ ID NO 132: from 1 to
35	<ul><li>(B) Polypeptide Sequence</li><li>- Pat. Appln. SEQ ID NO 133</li><li>- Ceres seq_id 1025686</li><li>- Location of start within SEQ ID NO 130: at 96 nt.</li></ul>
40	<ul> <li>(C) Nomination and Annotation of Domains within</li> <li>Predicted Polypeptide(s)         <ul> <li>Ribosomal protein L6</li> <li>Location within SEQ ID NO 133: from 3 to 185 aa.</li> </ul> </li> </ul>
45	(Dp) Related Amino Acid Sequences - Alignment No. 85 - gi No. 266945 - Description:
50	- % Identity: 84 - Alignment Length: 194

152

- Location of Alignment in SEQ ID NO 133: from 1 to 185

Maximum Length Sequence:

5 related to:

Clone IDs:

19274

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 134
- 10 Ceres seq id 1027152
  - Alternative transcription start site(s) located in SEQ ID NO 134:
    - -350, 3, 4, 10, 11, 13, 222
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 135
  - Ceres seq id 1027153
  - Location of start within SEQ ID NO 134: at 3 nt.
- 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 86
    - gi No. 2879811
- 25 Description:
  - % Identity: 86.6
  - Alignment Length: 112
  - Location of Alignment in SIO ID NO 135: from 26

to 137

30

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 136
  - Ceres seq id 1027154
  - Location of start within SEQ ID NO 134: at 78 nt.

35

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 87
    - qi No. 2879811
    - Description:
    - % Identity: 86.6
    - Alignment Length: 112
    - Location of Alignment in SEQ ID NO 136: from 1 to
- 45 112
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 137
    - Ceres seq id 1027155
- Location of start within SEQ ID NO 134: at 141 nt.

153

(C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 88 5 - qi No. 2879811 - Description: - % Identity: 86.6 - Alignment Length: 112 - Location of Alignment in SEQ ID NO 137: from 1 to 10 91 (Ba) Polypeptide Activities: Similar to ribosomal L30 protein activities. 15 Maximum Length Sequence: related to: Clone IDs: 17835 20 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 138 - Ceres seq id 1028095 - Alternative transcription start site(s) located in SEQ ID NO 138: 25 -2, 2, 3, 4, 5, 6, 12, 14, 18, 22, 26, 40, 42, 44, 45, 46, 47 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 139 - Ceres seq id 1028096 - Location of start within SEQ ID NO 138: at 55 nt. 30 - Location of Signal Peptide Cleavage Site within SEQ ID NO 139: at 29 aa. (C) Nomination and Annotation of Domains within 3.5 Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 89 - gi No. 4336325 - Description: 40 - % Identity: 31.7 - Alignment Length: 126 - Location of Alignment in SEQ ID NO 139: from 15 to 135 45 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 140 - Ceres seq id 1028097 - Location of start within SEQ ID NO 138: at 214 nt.

(C) Nomination and Annotation of Domains within

50

Predicted Polypeptide(s)

154

- (Dp) Related Amino Acid Sequences
  - Alignment No. 90
  - gi No. 4336325
  - Description:
  - % Identity: 31.7
  - Alignment Length: 126
  - Location of Alignment in SEQ ID NO 140: from 1 to

82

5

- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 141
  - Ceres seq id 1028098
  - Location of start within SEQ ID NO 138: at 3 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 15 ID NO 141: at 16 aa.
  - (Ba) Polypeptide Activities: Similar to human C214 membrane protein activities.

20

40

Maximum Length Sequence:

related to:

Clone IDs:

- 25 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 142
  - Ceres seq id 1028608
  - Alternative transcription start site(s) located in SEQ ID NO 142:
- 30 -4,2,28,31,36,49,59
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 143
    - Ceres seq id 1028609
- Location of start within SEQ ID NO 142: at 95 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 91
      - qi No. 2735528
      - Description:
      - % Identity: 33.9
      - Alignment Length: 118
- Location of Alignment in SEQ ID NO 143: from 64 to 178
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 144
- 50 Ceres seq id 1028610
  - Location of start within SEQ ID NO 142: at 176 nt.

	(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
5	(Dp) Related Amino Acid Sequences - Alignment No. 92 - gi No. 2735528 - Description:
L O	- % Identity: 33.9 - Alignment Length: 118 - Location of Alignment in SEQ ID NO 144: from 37 to 151
15	<ul> <li>(B) Polypeptide Sequence</li> <li>Pat. Appln. SEQ ID NO 145</li> <li>Ceres seq_id 1028611</li> <li>Location of start within SEQ ID NO 142: at 381 nt.</li> <li>Location of Signal Peptide Cleavage Site within SEQ</li> <li>ID NO 145: at 41 aa.</li> </ul>
20	<pre>(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)   (Dp) Related Amino Acid Sequences</pre>
25	Maximum Length Sequence: related to: Clone IDs: 1505
30	(Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 146 - Ceres seq_id 1030069
35	<ul> <li>(B) Polypeptide Sequence</li> <li>- Pat. Appln. SEQ ID NO 147</li> <li>- Ceres seq_id 1030070</li> <li>- Location of start within SEQ ID NO 146: at 2 nt.</li> </ul>
40	(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  (Dp) Related Amino Acid Sequences  - Alignment No. 93  - gi No. 4335755  - Description:
45	- % Identity: 63.6 - Alignment Length: 143 - Location of Alignment in SEQ ID NO 147: from 34 to 171
50	<ul><li>(B) Polypeptide Sequence</li><li>- Pat. Appln. SEQ ID NO 148</li><li>- Ceres seq_id 1030071</li></ul>

156

- Location of start within SEQ ID NO 146: at 50 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 94
    - qi No. 4335755
    - Description:
    - % Identity: 63.6
- Alignment Length: 143
  - Location of Alignment in SEQ ID NO 148: from 18 to 155
    - (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 149
    - Ceres seq id 1030072
    - Location of start within SEQ ID NO 146: at 170 nt.
- (C) Nomination and Annotation of Domains within 20 Predicted Polypeptide(s)
- (Dp) Related Amino Acid Sequences
  - Alignment No. 95
  - gi No. 4335755
  - Description:
- 25 % Identity: 63.6
  - Alignment Length: 143
  - Location of Alignment in SEQ ID NO 149: from 1 to 115
- 30 (Ba) Polypeptide Activities: Similar to hydroxyproline-rich protein activities.

Maximum Length Sequence:

35 related to:

5

15

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 150
- 40 Ceres seq id 1032069
  - Alternative transcription start site(s) located in SEQ ID NO 150:
    - 4,17
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 151
  - Ceres seq id 1032070
  - Location of start within SEQ ID NO 150: at 74 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

```
157
           (Dp) Related Amino Acid Sequences
               - Alignment No. 96
               - qi No. 3386621
               - Description:
5
               - % Identity:
                              97.4
               - Alignment Length: 234
               - Location of Alignment in SEQ ID NO 151: from 1 to
     231
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 152
            - Ceres seq id 1032071
            - Location of start within SEQ ID NO 150: at 122 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 97
               - qi No. 3386621
20
               - Description:
               - % Identity: 97.4
               - Alignment Length: 234
               - Location of Alignment in SEQ ID NO 152: from 1 to
     215
25
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
     Maximum Length Sequence:
30
      related to:
     Clone IDs:
           11466
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 153
35
          - Ceres seq id 1033557
          - Alternative transcription start site(s) located in SEQ
     ID NO 153:
            62, 64, 65, 67, 72, 73, 74, 75, 166
40
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 154
            - Ceres seq id 1033558
            - Location of start within SEQ ID NO 153: at 94 nt.
45
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein L34
               - Location within SEQ ID NO 154: from 105 to 145
     aa.
50
```

(Dp) Related Amino Acid Sequences

WO 00/40695 PCT/US00/00466 158 - Alignment No. 98 - qi No. 132909 - Description: - % Identity: 70 5 - Alignment Length: 30 - Location of Alignment in SEQ ID NO 154: from 116 to 145 (Ba) Polypeptide Activities: Similar to 50S ribosomal protein L34 activities. 10 Maximum Length Sequence: related to: Clone IDs: 15 21589 106951 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 155 - Ceres seq id 1034688 20 - Alternative transcription start site(s) located in SEQ ID NO 155: - Clone 21589 starts at 2 and ends at in cDNA. 25 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 156 - Ceres seq id 1034689 - Location of start within SEQ ID NO 155: at 2 nt. 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 99 - qi No. 4335755 35 - Description: - % Identity: 65 - Alignment Length: 143 - Location of Alignment in SEQ ID NO 156: from 33 to 170 40 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 157 - Ceres seq id 1034690 - Location of start within SEQ ID NO 155: at 47 nt. 45 (C) Nomination and Annotation of Domains within

Predicted Polypeptide(s)

50

(Dp) Related Amino Acid Sequences

- Alignment No. 100

- gi No. 4335755 - Description:

159

- % Identity: 65
- Alignment Length: 143
- Location of Alignment in SEQ ID NO 157: from 18

to 155

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 158
  - Ceres seq id 1034691
  - Location of start within SEQ ID NO 155: at 167 nt.

10

15

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 101
- gi No. 4335755
  - Description:
  - % Identity: 65
  - Alignment Length: 143
  - Location of Alignment in SEQ ID NO 158: from 1 to

20 115

(Ba) Polypeptide Activities: Similar to hydroxy proline rich glycoprotein activities.

25

40

Maximum Length Sequence:

related to:

Clone IDs:

10433

- 30 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 159
  - Ceres seg id 1035033
  - Alternative transcription start site(s) located in SEQ ID NO 159:

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 160
  - Ceres seq id 1035034
  - Location of start within SEQ ID NO 159: at 3 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 160: at 47 aa.
  - (C) Nomination and Annotation of Domains within
- 45 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 102
    - qi No. 3062795
    - Description:
- % Identity: 39.8
  - Alignment Length: 83

160

- Location of Alignment in SEQ ID NO 160: from 24 to 99  $\,$ 

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 161
  - Ceres seq id 1035035
  - Location of start within SEQ ID NO 159: at 69 nt.
- Location of Signal Peptide Cleavage Site within SEQ

ID NO 161: at 25 aa.

10

5

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 103
- gi No. 3062795
  - Description:
  - % Identity: 39.8
  - Alignment Length: 83
  - Location of Alignment in SEQ ID NO 161: from 2 to

20 77

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 162
  - Ceres seq id 1035036
- 25 Location of start within SEQ ID NO 159: at 72 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 162: at 24 aa.
- (C) Nomination and Annotation of Domains within 30 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 104
    - qi No. 3062795
    - Description:
    - % Identity: 39.8
    - Alignment Length: 83
    - Location of Alignment in SEQ ID NO 162: from 1 to

76

35

40 (Ba) Polypeptide Activities: Similar to Pollen coat protein activities.

Maximum Length Sequence:

45 related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 163
- 50 Ceres seq id 1035071

161

- Alternative transcription start site(s) located in SEO ID NO 163:

-2, -1, 2, 3, 18, 19, 21, 24, 25, 56, 66

- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 164
  - Ceres seq id 1035072
  - Location of start within SEQ ID NO 163: at 66 nt.
- 10 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - KH domain
  - Location within SEQ ID NO 164: from 47 to 95 aa.
- 15 (Dp) Related Amino Acid Sequences
  - Alignment No. 105
  - qi No. 133940
  - Description:
  - % Identity: 76
  - Alignment Length: 246
    - Location of Alignment in SEQ ID NO 164: from 1 to
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 165
    - Ceres seq id 1035073
    - Location of start within SEQ ID NO 163: at 195 nt.
- (C) Nomination and Annotation of Domains within 30
- Predicted Polypeptide(s)

20

25

35

- KH domain
- Location within SEQ ID NO 165: from 4 to 52 aa.
- (Dp) Related Amino Acid Sequences
- Alignment No. 106
  - gi No. 133940
  - Description:
  - % Identity: 76
  - Alignment Length: 246
- Location of Alignment in SEQ ID NO 165: from 1 to 40 105
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 166
- 45 - Ceres seq id 1035074
  - Location of start within SEQ ID NO 163: at 513 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 50 (Dp) Related Amino Acid Sequences
  - Alignment No. 107

162 - gi No. 133940 - Description: - % Identity: - Alignment Length: 246 - Location of Alignment in SEQ ID NO 166: from 1 to 5 97 Maximum Length Sequence: related to: 10 Clone IDs: 10511 Public Genomic DNA: qi No: 4539290 Predicted Exons: 15 INTR 5588 ... 5314 OCKHAMG-CDNA INTR 5517 ... 5314 OCKHAMG-CDNA qi No: 4914454 Predicted Exons: 20 38937 ... INTR 38663 OCKHAMG-CDNA 38866 ... 38663 INTR OCKHAMG-CDNA (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 167 25 - Ceres seq id 1376587 - Alternative transcription start site(s) located in SEQ ID NO 167: 8, 9, 10, 12, 13, 14, 15, 16, 17, 30, 34, 39, 41 30 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 168 - Ceres seq id 1376588 - Location of start within SEQ ID NO 167: at 2 nt. 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 108 - gi No. 4539292 - Description: 40 - % Identity: 99.4 - Alignment Length: 177 - Location of Alignment in SEQ ID NO 168: from 26 to 202 45 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 169 - Ceres seq\_id 1376589

- Location of start within SEQ ID NO 167: at 77 nt.

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 109
               - qi No. 4539292
 5
               - Description:
               - % Identity:
                              99.4
               - Alignment Length: 177
               - Location of Alignment in SEQ ID NO 169: from 1 to
     177
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 170
            - Ceres seq id 1376590
15
            - Location of start within SEQ ID NO 167: at 212 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
20
               - Alignment No. 110
               - qi No. 4539292
               - Description:
               - % Identity: 99.4
               - Alignment Length: 177
25
               - Location of Alignment in SEQ ID NO 170: from 1 to
     132
     (Ba) Polypeptide Activities: Similar 40S ribosomal protein
     activities.
30
     Maximum Length Sequence:
      related to:
     Clone IDs:
           112110
35
     Public Genomic DNA:
           gi No: 4263774
           Predicted Exons:
                     1816
              INIT
                               1814
                                         OCKHAMG-CDS
              INTR
                     1290 ... 1112
                                         OCKHAMG-CDS
40
              TERM
                     959
                            ... 803
                                         OCKHAMG-CDS
           gi No: 4510360
           Predicted Exons:
                     114707... 114705 OCKHAMG-CDS
              INIT
                     114181... 114003 OCKHAMG-CDS
              INTR
45
              TERM 113850... 113694
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 171
          - Ceres seq id 1378581
          - Alternative transcription start site(s) located in SEQ
50
     ID NO 171:
            -35, -4, -3, 18, 20
```

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(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 172
            - Ceres seq id 1378582
5
            - Location of start within SEQ ID NO 171: at 112 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 172: at 17 aa.
           (C) Nomination and Annotation of Domains within
10
     Predicted Polypeptide(s)
               - DnaJ domain
               - Location within SEQ ID NO 172: from 57 to 101 aa.
           (Dp) Related Amino Acid Sequences
15
               - Alignment No. 111
               - qi No. 4263775
               - Description:
               - % Identity:
                              100
               - Alignment Length: 112
20
               - Location of Alignment in SEQ ID NO 172: from 1 to
     112
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 173
25
            - Ceres seq id 1378583
            - Location of start within SEQ ID NO 171: at 256 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
30
               - DnaJ domain
               - Location within SEQ ID NO 173: from 9 to 53 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 112
               - qi No. 4263775
35
               - Description:
               - % Identity: 100
               - Alignment Length: 112
               - Location of Alignment in SEQ ID NO 173: from 1 to
40
     64
     Maximum Length Sequence:
      related to:
     Clone IDs:
45
           13599
      (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 174
          - Ceres seq id 1383462
          - Alternative transcription start site(s) located in SEQ
50
     ID NO 174:
            -2,2,3,4,5,6,8,14,18,24
```

50

165

(B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 175 - Ceres seq id 1383463 - Location of start within SEQ ID NO 174: at 3 nt. 5 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Plant lipid transfer protein family 10 - Location within SEQ ID NO 175: from 46 to 119 aa. (Dp) Related Amino Acid Sequences - Alignment No. 113 - qi No. 3128176 15 - Description: - % Identity: 37 - Alignment Length: 194 - Location of Alignment in SEQ ID NO 175: from 23 to 197 20 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 176 - Ceres seq id 1383464 - Location of start within SEQ ID NO 174: at 42 nt. 25 - Location of Signal Peptide Cleavage Site within SEQ ID NO 176: at 24 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Plant lipid transfer protein family 30 - Location within SEQ ID NO 176: from 33 to 106 aa. (Dp) Related Amino Acid Sequences - Alignment No. 114 35 - gi No. 3128176 - Description: - % Identity: 37 - Alignment Length: 194 - Location of Alignment in SEQ ID NO 176: from 10 40 to 184 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 177 - Ceres seq id 1383465 45 - Location of start within SEQ ID NO 174: at 90 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Plant lipid transfer protein family

- Location within SEQ ID NO 177: from 17 to 90 aa.

WO 00/40695 PCT/US00/00466 166 (Dp) Related Amino Acid Sequences - Alignment No. 115 - gi No. 3128176 - Description: - % Identity: 5 37 - Alignment Length: 194 - Location of Alignment in SEQ ID NO 177: from 1 to 168 10 Maximum Length Sequence: related to: Clone IDs: 156375 (Ac) cDNA Polynucleotide Sequence 15 - Pat. Appln. SEQ ID NO 178 - Ceres seq id 1386215 - Alternative transcription start site(s) located in SEQ ID NO 178: -38, 12, 17, 18, 19, 20, 26 20 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 179 - Ceres seq\_id 1386216 - Location of start within SEQ ID NO 178: at 2 nt. 25 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Photosystem I psaG / psaK - Location within SEQ ID NO 179: from 77 to 138 aa. 30 (Dp) Related Amino Acid Sequences - Alignment No. 116 - gi No. 3885511 - Description: 35 - % Identity: 79.3 - Alignment Length: 112 - Location of Alignment in SEQ ID NO 179: from 28 to 138 40 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 180 - Ceres seq id 1386217 - Location of start within SEQ ID NO 178: at 83 nt. 45 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Photosystem I psaG / psaK - Location within SEQ ID NO 180: from 50 to 111 aa.

(Dp) Related Amino Acid Sequences

- Alignment No. 117

167

```
- qi No. 3885511
```

- Description:
- % Identity: 79.3
- Alignment Length: 112
- 5 Location of Alignment in SEQ ID NO 180: from 1 to 111
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 181
- Ceres seq id 1386218
  - Location of start within SEQ ID NO 178: at 95 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Photosystem I psaG / psaK
    - Location within SEQ ID NO 181: from 46 to 107 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 118
      - qi No. 3885511
      - Description:
      - % Identity: 79.3
      - Alignment Length: 112
      - Location of Alignment in SEQ ID NO 181: from 1 to
- 25 107

15

20

35

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 182
  - Ceres seq id 2025156
- Location of start within SEQ ID NO 178: at 502 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
    - (Dp) Related Amino Acid Sequences

Maximum Length Sequence:

related to:

Clone IDs:

- 40 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 183
  - Ceres seq id 1388499
  - Alternative transcription start site(s) located in SEQ ID NO 183:
- 45 2,5,6,10,16,30,89,346,349
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 184
    - Ceres seq id 1388500
- 50 Location of start within SEQ ID NO 183: at 62 nt.

168

(C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 119 5 - qi No. 2829899 - Description: - % Identity: 49.3 - Alignment Length: 150 - Location of Alignment in SEQ ID NO 184: from 2 to 10 150 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 185 - Ceres seq id 1388501 15 - Location of start within SEQ ID NO 183: at 122 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 20 - Alignment No. 120 - gi No. 2829899 - Description: - % Identity: 49.3 - Alignment Length: 150 - Location of Alignment in SEQ ID NO 185: from 1 to 25 130 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 186 - Ceres seq id 1388502 30 - Location of start within SEQ ID NO 183: at 266 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) 35 (Dp) Related Amino Acid Sequences - Alignment No. 121 - qi No. 2829899 - Description: - % Identity: 49.3 - Alignment Length: 150 40 - Location of Alignment in SEQ ID NO 186: from 1 to 82 (Ba) Polypeptide Activities: Similar to major latex protein 45 activities.

Maximum Length Sequence:

50 related to: Clone IDs:

169

21304

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 187
  - Ceres seq id 1388519

5

20

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 188
  - Ceres seq id 1388520
- 10 Location of start within SEQ ID NO 187: at 1 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 188: at 38 aa.
  - (C) Nomination and Annotation of Domains within
- 15 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 122
    - qi No. 4584110
    - Description:
    - % Identity: 48.5
    - Alignment Length: 163
    - Location of Alignment in SEQ ID NO 188: from 20
  - to 182
- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 189
  - Ceres seq id 1388521
  - Location of start within SEQ ID NO 187: at 58 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 30 ID NO 189: at 19 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 123
        - qi No. 4584110
        - Description:
        - % Identity: 48.5
        - Alignment Length: 163
- Location of Alignment in SEQ ID NO 189: from 1 to 163
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 190
- 45 Ceres seq id 1388522
  - Location of start within SEQ ID NO 187: at 73 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 190: at 14 aa.
- 50 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

170 (Dp) Related Amino Acid Sequences - Alignment No. 124 - qi No. 4584110 - Description: - % Identity: 5 48.5 - Alignment Length: 163 - Location of Alignment in SEQ ID NO 190: from 1 to 158 10 (Ba) Polypeptide Activities: Similar to pollen specific protein activities. Maximum Length Sequence: related to: 15 Clone IDs: 2153 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 191 - Ceres seq id 1388563 20 - Alternative transcription start site(s) located in SEQ ID NO 191: -3, -2, -1, 13, 15, 146(B) Polypeptide Sequence 25 - Pat. Appln. SEQ ID NO 192 - Ceres seq id 1388564 - Location of start within SEQ ID NO 191: at 2 nt. (C) Nomination and Annotation of Domains within 30 Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 193 35 - Ceres seq id 1388565 - Location of start within SEQ ID NO 191: at 306 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) 40 (Dp) Related Amino Acid Sequences - Alignment No. 125 - qi No. 3927834 - Description: - % Identity: 31.5 - Alignment Length: 108 45 - Location of Alignment in SEQ ID NO 193: from 1 to

(B) Polypeptide Sequence

43

50

- Pat. Appln. SEQ ID NO 194

- Ceres seq id 1388566

171

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- Location of start within SEQ ID NO 191: at 455 nt.
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- Location of Signal Peptide Cleavage Site within SEQ ID NO 194: at 36 aa.

5 (Ba) Polypeptide Activities: Similar to hydroxproline-rich glycoprotein activities.

Maximum Length Sequence:

10 related to:

25

45

Clone IDs:

22488

Public Genomic DNA:

qi No: 5708384

15 Predicted Exons:

INIT 82228 ... 82063 OCKHAMG-CDS TERM 81730 ... 81381 OCKHAMG-CDS

gi No: 5732090

Predicted Exons:

20 INIT 21301 ... 21136 OCKHAMG-CDS TERM 20803 ... 20454 OCKHAMG-CDS

gi No: 5870169

Predicted Exons:

INIT 89258 ... 89423 OCKHAMG-CDS TERM 89756 ... 90105 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 195
- Ceres seq id 1388793
- Alternative transcription start site(s) located in SEQ 30 ID NO 195:
  - 7,14,15,20,24,30,33,42,44,58,63,64,66,76,94,97,98,99,101,102
- 35 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 196
  - Ceres seq id 1388794
  - Location of start within SEQ ID NO 195: at 154 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 40 ID NO 196: at 19 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
    - Alignment No. 126
      - gi No. 4584110
      - Description:
      - % Identity: 48.5
      - Alignment Length: 163
- Location of Alignment in SEQ ID NO 196: from 1 to

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 197
            - Ceres seq id 1388795
 5
            - Location of start within SEQ ID NO 195: at 169 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 197: at 14 aa.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
10
           (Dp) Related Amino Acid Sequences
               - Alignment No. 127
               - gi No. 4584110
               - Description:
15
               - % Identity:
                             48.5
               - Alignment Length: 163
               - Location of Alignment in SEQ ID NO 197: from 1 to
     158
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 198
            - Ceres seq id 1388796
            - Location of start within SEQ ID NO 195: at 205 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 128
               - qi No. 4584110
               - Description:
30
               - % Identity: 48.5
               - Alignment Length: 163
               - Location of Alignment in SEQ ID NO 198: from 1 to
     146
35
     (Ba) Polypeptide Activities: Similar to pollen specific
     protein activities.
     Maximum Length Sequence:
40
      related to:
     Clone IDs:
           26569
     Public Genomic DNA:
           qi No: 6143856
45
           Predicted Exons:
                     78628 ... 78255 OCKHAMG-CDNA
              INTR
                     78160 ... 78004
              INTR
                                        OCKHAMG-CDNA
              INTR
                     78624 ... 78255 OCKHAMG-CDNA
50
              INTR 78160 ... 78004
                                        OCKHAMG-CDNA
```

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173
                    78624 ...
                                78255 OCKHAMG-CDNA
              INTR
                                78006 OCKHAMG-CDNA
                    78160 ...
              INTR
                    78624 ...
                               78283
              INTR
                                       OCKHAMG-CDNA
 5
           qi No: 6223633
           Predicted Exons:
              INTR
                    78629 ...
                               78256
                                        OCKHAMG-CDNA
                    78161 ...
              INTR
                               78005
                                        OCKHAMG-CDNA
                    78625 ...
10
              INTR
                               78256
                                        OCKHAMG-CDNA
                    78161 ...
                               78005
              INTR
                                        OCKHAMG-CDNA
                    78625 ... 78256 OCKHAMG-CDNA
              INTR
              INTR
                    78161 ...
                               78007
                                        OCKHAMG-CDNA
15
                    78625 ...
                               78284 OCKHAMG-CDNA
              INTR
                     78587 ... 78256
              INIT
                                       OCKHAMG-CDS
              TERM
                     78161 ... 78152
                                        OCKHAMG-CDS
20
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 199
          - Ceres seq id 1392041
          - Alternative transcription start site(s) located in SEQ
     ID NO 199:
25
            3, 4, 5, 6, 7, 12
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 200
            - Ceres seq id 1392042
30
            - Location of start within SEQ ID NO 199: at 45 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 200: at 19 aa.
           (C) Nomination and Annotation of Domains within
35
     Predicted Polypeptide(s)
               - Plant lipid transfer protein family
               - Location within SEQ ID NO 200: from 23 to 112 aa.
           (Dp) Related Amino Acid Sequences
40
               - Alignment No. 129
               - qi No. 2497753
               - Description:
               - % Identity:
                             47
               - Alignment Length: 117
45
               - Location of Alignment in SEQ ID NO 200: from 4 to
     113
     Maximum Length Sequence:
```

related to:
50 Clone IDs:

WO 00/40095 FC1/0500/0040

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174
    Public Genomic DNA:
          gi No: 5731404
          Predicted Exons:
                    70440 ...
             INTR
                               70346
                                       OCKHAMG-CDNA
                    70076 ...
5
             INTR
                               69865
                                       OCKHAMG-CDNA
                    69508 ...
                               69434 OCKHAMG-CDNA
             INTR
             INTR
                    69358 ... 69284 OCKHAMG-CDNA
             INTR 69195 ... 69115 OCKHAMG-CDNA
             INTR 69033 ... 68959 OCKHAMG-CDNA
10
             INTR 68868 ... 68695 OCKHAMG-CDNA
             INTR 68581 ... 68344 OCKHAMG-CDNA
                    70056 ... 69865
             INIT
                                      OCKHAMG-CDS
             INTR
                    69508 ... 69434 OCKHAMG-CDS
                    69358 ... 69284 OCKHAMG-CDS
15
             INTR
                    69195 ... 69115 OCKHAMG-CDS
             INTR
             INTR
                    69033 ... 68959 OCKHAMG-CDS
                    68868 ... 68695 OCKHAMG-CDS
             INTR
             TERM
                    68581 ... 68546 OCKHAMG-CDS
20
     (Ac) cDNA Polynucleotide Sequence
         - Pat. Appln. SEQ ID NO 201
         - Ceres seq id 1393342
         - Alternative transcription start site(s) located in SEQ
     ID NO 201:
25
            16,48
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 202
            - Ceres seq id 1393343
30
            - Location of start within SEQ ID NO 201: at 118 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
35
               - Alignment No. 130
              - qi No. 3386621
               - Description:
              - % Identity:
               - Alignment Length: 224
40
               - Location of Alignment in SEQ ID NO 202: from 2 to
     225
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
45
     Maximum Length Sequence:
      related to:
     Clone IDs:
           29453
50
     (Ac) cDNA Polynucleotide Sequence
```

- Pat. Appln. SEQ ID NO 203

175

- Ceres seq id 1393556

- Alternative transcription start site(s) located in SEQ ID NO 203:

7,11,14,23,37,41,42,57,61

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 204
  - Ceres seq id 1393557
  - Location of start within SEQ ID NO 203: at 3 nt.

10

15

- - (Dp) Related Amino Acid Sequences
    - Alignment No. 131
  - qi No. 1082054
    - Description:
    - % Identity: 36.1
    - Alignment Length: 61
    - Location of Alignment in SEQ ID NO 204: from 23

20 to 82

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 205
  - Ceres seq id 1393558
- 25 Location of start within SEQ ID NO 203: at 69 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 205: at 41 aa.
- (C) Nomination and Annotation of Domains within 30 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 132
    - qi No. 1082054
    - Description:
    - % Identity: 36.
    - Alignment Length: 61
    - Location of Alignment in SEQ ID NO 205: from 1 to

60

35

- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 206
  - Ceres seq id 1393559
  - Location of start within SEQ ID NO 203: at 72 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 45 ID NO 206: at 40 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 133
      - gi No. 1082054

176

- Description:
- % Identity: 36.1
- Alignment Length: 61
- Location of Alignment in SEQ ID NO 206: from 1 to

5 59

(Ba) Polypeptide Activities: Similar to transmembrane copper transporter protein activities.

10

30

40

Maximum Length Sequence:

related to:

Clone IDs:

- 15 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 207
  - Ceres seq id 1396782
  - Alternative transcription start site(s) located in SEQ ID NO 207:
- 20 -1,29
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 208
    - Ceres seq id 1396783
- 25 Location of start within SEQ ID NO 207: at 3 nt.
  - - (Dp) Related Amino Acid Sequences
    - Alignment No. 134
      - qi No. 4512613
      - Description:
      - % Identity: 99.2
      - Alignment Length: 122
- Location of Alignment in SEQ ID NO 208: from 11 to 132
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 209
  - Ceres seq id 1396784
    - Location of start within SEQ ID NO 207: at 33 nt.
    - Location of Signal Peptide Cleavage Site within SEQ ID NO 209: at 19 aa.
- 45 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 135
    - gi No. 4512613
- 50 Description:
  - % Identity: 99.2

177

- Alignment Length: 122

- Location of Alignment in SEQ ID NO 209: from 1 to

122

- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 210
  - Ceres seq id 1396785
  - Location of start within SEQ ID NO 207: at 45 nt.
- 10 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 136
    - gi No. 4512613
- 15 Description:
  - % Identity: 99.2
  - Alignment Length: 122
  - Location of Alignment in SEQ ID NO 210: from 1 to

118

20

(Ba) Polypeptide Activities: Similar to intergenic region of cyt2-MDH1 of yeast activities.

Maximum Length Sequence:

25 related to:

Clone IDs:

42402

- (Ac) cDNA Polynucleotid€ Sequence
  - Pat. Appln. SEQ ID NO 211
- 30 Ceres seq id 1396802
  - Alternative transcription start site(s) located in SEQ ID NO 211:

-145,-56

- 35 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 212
  - Ceres seq\_id 1396803
  - Location of start within SEQ ID NO 211: at 1 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 40 ID NO 212: at 19 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
- 45 Alignment No. 137
  - gi No. 2463339
  - Description:
  - % Identity: 38.6
  - Alignment Length: 70
- Location of Alignment in SEQ ID NO 212: from 136 to 205

178

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 213
            - Ceres seq id 1396804
 5
            - Location of start within SEQ ID NO 211: at 136 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 138
10
               - gi No. 2463339
               - Description:
               - % Identity:
                              38.6
               - Alignment Length: 70
15
               - Location of Alignment in SEQ ID NO 213: from 91
     to 160
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 214
20
            - Ceres seq id 1396805
            - Location of start within SEO ID NO 211: at 172 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
25
               - Alignment No. 139
               - qi No. 2463339
               - Description:
               - % Identity:
                              38.6
30
               - Alignment Length: 70
               - Location of Alignment in SEQ ID NO 214: from 79
     to 148
     (Ba) Polypeptide Activities: Similar to rbcX protein
35
     activities.
     Maximum Length Sequence:
      related to:
     Clone IDs:
40
           5105
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 215
          - Ceres seq id 1397130
          - Alternative transcription start site(s) located in SEQ
45
     ID NO 215:
            -1
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 216
            - Ceres seq id 1397131
50
```

- Location of start within SEQ ID NO 215: at 1 nt.

179

5	<pre>(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)   (Dp) Related Amino Acid Sequences</pre>
15	<ul> <li>(B) Polypeptide Sequence</li> <li>- Pat. Appln. SEQ ID NO 217</li> <li>- Ceres seq_id 1397132</li> <li>- Location of start within SEQ ID NO 215: at 52 nt.</li> </ul>
20	<ul> <li>(C) Nomination and Annotation of Domains within</li> <li>Predicted Polypeptide(s)</li> <li>(Dp) Related Amino Acid Sequences</li> <li>Alignment No. 141</li> <li>gi No. 2827551</li> </ul>
25	<ul> <li>Description:</li> <li>% Identity: 63.1</li> <li>Alignment Length: 179</li> <li>Location of Alignment in SEQ ID NO 217: from 1 to</li> </ul>
30	<ul><li>(B) Polypeptide Sequence</li><li>- Pat. Appln. SEQ ID NO 218</li><li>- Ceres seq_id 1397133</li><li>- Location of start within SEQ ID NO 215: at 211 nt.</li></ul>
35	(C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 142
40	<pre>- gi No. 2827551 - Description: - % Identity: 63.1 - Alignment Length: 179 - Location of Alignment in SEQ ID NO 218: from 1 to 106</pre>
45	(Ba) Polypeptide Activities: Similar to meth CpG binding protein activities.
50	Maximum Length Sequence: related to:

Clone IDs:

180

92204

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 219
- Ceres seq id 1398004
- 5 Alternative transcription start site(s) located in SEQ ID NO 219:

2,3,5,8,28

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 220
  - Ceres seq id 1398005
  - Location of start within SEQ ID NO 219: at 68 nt.
- (C) Nomination and Annotation of Domains within
- 15 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 143
    - gi No. 4454037
    - Description:
    - % Identity: 99.3
    - Alignment Length: 151
  - Location of Alignment in SEQ ID NO 220: from 1 to 151
- (Ba) Polypeptide Activities: Similar to major latex protein activities.

Maximum Length Sequence:

related to:

Clone IDs:

98584

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 221
  - Ceres seq id 1399370

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30

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 222
  - Ceres seq id 1399371
  - Location of start within SEQ ID NO 221: at 3 nt.

40

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 144
    - gi No. 4886285
    - Description:
    - % Identity: 28.8
    - Alignment Length: 59
    - Location of Alignment in SEQ ID NO 222: from 25
- 50 to 83

181

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(B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 223
- Ceres seq id 1399372
- Location of start within SEQ ID NO 221: at 45 nt.

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30

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 145
    - gi No. 4886285
      - Description:
      - % Identity: 28.8
      - Alignment Length: 59
    - Location of Alignment in SEQ ID NO 223: from 11
- 15 to 69
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 224
    - Ceres seq id 1399373
- 20 Location of start within SEQ ID NO 221: at 60 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 224: at 25 aa.
- (C) Nomination and Annotation of Domains within
- 25 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 140
    - gi No. 4886285
    - Description:
  - % Identity: 28.8
    - Alignment Length: 59
    - Location of Alignment in SEQ ID NO 224: from 6 to 64
- 35 (Ba) Polypeptide Activities: Similar to outer envelope membrane protein in choloroplast in pea activities.

Maximum Length Sequence:

40 related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 225
- 45 Ceres seq id 1425147
  - Alternative transcription start site(s) located in SEQ ID NO 225:
    - 20,31,49,58,80
- 50 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 226

182

- Ceres seq id 1425148
- Location of start within SEQ ID NO 225: at 85 nt.
- (C) Nomination and Annotation of Domains within
  Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 147
    - qi No. 3510256
    - Description:
- % Identity: 37.8
  - Alignment Length: 158
  - Location of Alignment in SEQ ID NO 226: from 1 to 155
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 227
  - Ceres seq id 1425149
  - Location of start within SEQ ID NO 225: at 241 nt.
- 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 148
    - gi No. 3510256
- 25 Description:
  - % Identity: 37.8
  - Alignment Length: 158
  - Location of Alignment in SEQ ID NO 227: from 1 to

103

30

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

Maximum Length Sequence:

35 related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 228
- 40 Ceres seq id 1441102
  - Alternative transcription start site(s) located in SEQ ID NO 228:
    - 2, 3, 4, 5, 8, 10, 16, 22, 51, 69, 388
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 229
  - Ceres seq id 1441103
  - Location of start within SEQ ID NO 228: at 1 nt.
- 50 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

183

- (Dp) Related Amino Acid Sequences
  - Alignment No. 149
  - gi No. 4337175
  - Description:
  - % Identity: 100
  - Alignment Length: 179
  - Location of Alignment in SEQ ID NO 229: from 19

to 197

5

- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 230
  - Ceres seq id 1441104
  - Location of start within SEQ ID NO 228: at 55 nt.
- 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 150
    - gi No. 4337175
- 20 Description:
  - % Identity: 100
  - Alignment Length: 179
  - Location of Alignment in SEQ ID NO 230: from 1 to

179

25

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 231
  - Ceres seq\_id 1441105
  - Location of start within SEQ ID NO 228: at 94 nt.

30

35

- - (Dp) Related Amino Acid Sequences
    - Alignment No. 151
    - gi No. 4337175
    - Description:
    - % Identity: 100
    - Alignment Length: 179
    - Location of Alignment in SEQ ID NO 231: from 1 to

40 166

- (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.
- 45 Maximum Length Sequence:

related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
- 50 Pat. Appln. SEQ ID NO 232
  - Ceres seq id 1447480

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 233
5
            - Ceres seq id 1447481
            - Location of start within SEQ ID NO 232: at 52 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
10
           (Dp) Related Amino Acid Sequences
               - Alignment No. 152
               - qi No. 3510256
               - Description:
               - % Identity:
                              54
15
               - Alignment Length: 50
               - Location of Alignment in SEQ ID NO 233: from 77
    to 126
        (B) Polypeptide Sequence
20
            - Pat. Appln. SEQ ID NO 234
            - Ceres seq id 1447482
            - Location of start within SEQ ID NO 232: at 208 nt.
           (C) Nomination and Annotation of Domains within
25
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 153
               - gi No. 3510256
               - Description:
30
               - % Identity:
                              54
               - Alignment Length: 50
               - Location of Alignment in SEQ ID NO 234: from 25
     to 74
35
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
     Maximum Length Sequence:
      related to:
40
     Clone IDs:
           268712
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 235
          - Ceres seq id 1447577
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 236
            - Ceres seq id 1447578
50
            - Location of start within SEQ ID NO 235: at 2 nt.
```

```
185
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 154
 5
               - qi No. 5080769
               - Description:
               - % Identity:
                              100
               - Alignment Length: 17
               - Location of Alignment in SEQ ID NO 236: from 16
10
     to 32
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 237
            - Ceres seq id 1447579
15
            - Location of start within SEQ ID NO 235: at 165 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
20
               - Alignment No. 155
               - gi No. 5080769
               - Description:
               - % Identity:
                              91.5
               - Alignment Length: 153
25
               - Location of Alignment in SEQ ID NO 237: from 1 to
     115
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 238
30
            - Ceres seq id 1447580
            - Location of start within SEQ ID NO 235: at 298 nt.
           (Ba) Polypeptide Activities: Similar to
     hydroxyproline-rich glycoprotein activities.
35
     Maximum Length Sequence:
      related to:
     Clone IDs:
40
           94821
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 239
          - Ceres seq id 1447922
          - Alternative transcription start site(s) located in SEQ
45
     ID NO 239:
            12,16
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 240
```

- Location of start within SEQ ID NO 239: at 81 nt.

- Ceres seq id 1447923

186

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(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
5
               - Alignment No. 156
               - qi No. 3510256
               - Description:
               - % Identity:
                              37.8
               - Alignment Length: 158
10
               - Location of Alignment in SEQ ID NO 240: from 1 to
     155
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 241
            - Ceres seq id 1447924
15
           - Location of start within SEQ ID NO 239: at 237 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
20
           (Dp) Related Amino Acid Sequences
               - Alignment No. 157
               - gi No. 3510256
               - Description:
               - % Identity: 37.8
25
               - Alignment Length: 158
               - Location of Alignment in SEQ ID NO 241: from 1 to
     103
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
30
     dicot specific gene, plant specific gene.
     Maximum Length Sequence:
      related to:
     Clone IDs:
35
           20539
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 242
          - Ceres seq id 1448012
          - Alternative transcription start site(s) located in SEQ
40
     ID NO 242:
            -30.-6
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 243
45
            - Ceres seq id 1448013
            - Location of start within SEQ ID NO 242: at 79 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
50
               - Ribosomal protein S16
```

- Location within SEQ ID NO 243: from 9 to 70 aa.

187

```
(Dp) Related Amino Acid Sequences
               - Alignment No. 158
               - gi No. 3096931
 5
               - Description:
               - % Identity:
                               57.5
               - Alignment Length: 113
               - Location of Alignment in SEQ ID NO 243: from 1 to
     112
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 244
            - Ceres seq id 1448014
            - Location of start within SEQ ID NO 242: at 139 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein S16
               - Location within SEQ ID NO 244: from 1 to 50 aa.
20
           (Dp) Related Amino Acid Sequences
               - Alignment No. 159
               - qi No. 3096931
               - Description:
               - % Identity:
25
                               57.5
               - Alignment Length: 113
               - Location of Alignment in SEQ ID NO 244: from 1 to
     92
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 245
            - Ceres seq id 1448015
            - Location of start within SEQ ID NO 242: at 232 nt.
35
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
               - Alignment No. 160
                - gi No. 3096931
               - Description:
40
                - % Identity:
                               57.5
                - Alignment Length: 113
                - Location of Alignment in SEQ ID NO 245: from 1 to
     61
45
     Maximum Length Sequence:
      related to:
     Clone IDs:
           34091
50
     (Ac) cDNA Polynucleotide Sequence
```

- Pat. Appln. SEQ ID NO 246

```
188
          - Ceres seq id 1448135
          - Alternative transcription start site(s) located in SEQ
     ID NO 246:
            -1
5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 247
            - Ceres seq id 1448136
            - Location of start within SEQ ID NO 246: at 3 nt.
10
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
15
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 248
            - Ceres seq id 1448137
            - Location of start within SEQ ID NO 246: at 68 nt.
20
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein S14p/S29e
               - Location within SEQ ID NO 248: from 3 to 54 aa.
25
           (Dp) Related Amino Acid Sequences
               - Alignment No. 161
               - gi No. 4506717
               - Description:
               - % Identity: 72.2
30
               - Alignment Length: 54
               - Location of Alignment in SEQ ID NO 248: from 1 to
     54
        (B) Polypeptide Sequence
35
            - Pat. Appln. SEQ ID NO 249
            - Ceres seq id 1448138
            - Location of start within SEQ ID NO 246: at 85 nt.
           (C) Nomination and Annotation of Domains within
40
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
     Maximum Length Sequence:
      related to:
4.5
     Clone IDs:
           39285
     Public Genomic DNA:
           qi No: 4406776
           Predicted Exons:
```

SINGLE 61584 ... 61895 GENBANK

189 SINGLE 61584 ... 61895 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 250 - Ceres seq id 1448185 - Alternative transcription start site(s) located in SEQ 5 ID NO 250: -36, -18, -17, 16, 19 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 251 10 - Ceres seq id 1448186 - Location of start within SEQ ID NO 250: at 41 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) 15 (Dp) Related Amino Acid Sequences - Alignment No. 162 - gi No. 4406787 - Description: 20 - % Identity: 100 - Alignment Length: 103 - Location of Alignment in SEQ ID NO 251: from 1 to 103 25 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 252 - Ceres seq id 1448187 - Location of start within SEQ ID NO 250: at 68 nt. 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 163 - qi No. 4406787 35 - Description: - % Identity: 100 - Alignment Length: 103 - Location of Alignment in SEQ ID NO 252: from 1 to 94 40 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 253 - Ceres seq id 1448188 - Location of start within SEQ ID NO 250: at 89 nt. 45 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 164 50 - qi No. 4406787

- Description:

190

- % Identity: 100
- Alignment Length: 103
- Location of Alignment in SEQ ID NO 253: from 1 to 87
- 5 (Ba) Polypeptide Activities: Similar to NADH dehydrogenase protein activities
- 10 Maximum Length Sequence:

related to:

Clone IDs:

99119

- (Ac) cDNA Polynucleotide Sequence
- Pat. Appln. SEQ ID NO 254
  - Ceres seq id 1450875
  - (B) Polypeptide Sequence
- 20 Pat. Appln. SEQ ID NO 255
  - Ceres seq id 1450876
  - Location of start within SEQ ID NO 254: at 2 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 255: at 26 aa.

25

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- ATPases associated with various cellular activities (AAA)  $\,$
- Location within SEQ ID NO 255: from 391 to 598 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 165
    - qi No. 4309734
    - Description:
    - % Identity: 86
    - Alignment Length: 633
    - Location of Alignment in SEQ ID NO 255: from 9 to
- 40 634
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 256
  - Ceres seq id 1450877
- 45 Location of start within SEQ ID NO 254: at 14 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 256: at 22 aa.
- (C) Nomination and Annotation of Domains within 50 Predicted Polypeptide(s)

191 - ATPases associated with various cellular activities (AAA) - Location within SEQ ID NO 256: from 387 to 594 aa. 5 (Dp) Related Amino Acid Sequences - Alignment No. 166 - gi No. 4309734 - Description: 10 - % Identity: 86 - Alignment Length: 633 - Location of Alignment in SEQ ID NO 256: from 5 to 630 15 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 257 - Ceres seq id 1450878 - Location of start within SEQ ID NO 254: at 68 nt. 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - ATPases associated with various cellular activities (AAA) - Location within SEQ ID NO 257: from 369 to 576 2.5 aa. (Dp) Related Amino Acid Sequences - Alignment No. 167 - qi No. 4309734 30 - Description: - % Identity: 86 - Alignment Length: 633 - Location of Alignment in SEQ ID NO 257: from 1 to 612 35 Maximum Length Sequence: related to: Clone IDs: 269321 40 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 258 - Ceres seq id 1459191 (B) Polypeptide Sequence 45

- Pat. Appln. SEQ ID NO 259
- Ceres seq id 1459192
- Location of start within SEQ ID NO 258: at 1 nt.
- 50 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

```
192
           (Dp) Related Amino Acid Sequences
               - Alignment No. 168
               - gi No. 4490728
               - Description:
               - % Identity: 100
 5
               - Alignment Length: 17
               - Location of Alignment in SEQ ID NO 259: from 21
     to 37
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 260
            - Ceres seq id 1459193
            - Location of start within SEQ ID NO 258: at 3 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 169
               - qi No. 4490728
               - Description:
20
               - % Identity: 85.9
               - Alignment Length: 85
               - Location of Alignment in SEQ ID NO 260: from 37
     to 110
25
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
     Maximum Length Sequence:
30
      related to:
     Clone IDs:
           8446
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 261
35
          - Ceres seq id 1461848
          - Alternative transcription start site(s) located in SEQ
     ID NO 261:
            -3, 6, 7, 11, 31, 32, 34, 62, 63, 69, 70, 71, 97
40
         (B) Polypeptide Sequence
             - Pat. Appln. SEQ ID NO 262
             - Ceres seq id 1461849
             - Location of start within SEQ ID NO 261: at 113 nt.
             - Location of Signal Peptide Cleavage Site within SEQ
45
     ID NO 262: at 36 aa.
            (C) Nomination and Annotation of Domains within
```

- Predicted Polypeptide(s)
  - MAPEG family
- Location within SEQ ID NO 262: from 59 to 136 aa. 50

WO 00/40695 PCT/US00/00466 193 (Dp) Related Amino Acid Sequences - Alignment No. 170 - gi No. 4758714 - Description: 5 - % Identity: 41.9 - Alignment Length: 131 - Location of Alignment in SEQ ID NO 262: from 8 to 136 10 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 263 - Ceres seq id 1461850 - Location of start within SEQ ID NO 261: at 197 nt. 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - MAPEG family - Location within SEQ ID NO 263: from 31 to 108 aa. 20 (Dp) Related Amino Acid Sequences - Alignment No. 171 - qi No. 4758714 - Description: - % Identity: 41.9 25 - Alignment Length: 131 - Location of Alignment in SEQ ID NO 263: from 1 to 108 (B) Polypeptide Sequence 30 - Pat. Appln. SEQ ID NO 264 - Ceres seq id 1461851 - Location of start within SEQ ID NO 261: at 329 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 264: at 22 aa. 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - MAPEG family - Location within SEQ ID NO 264: from 1 to 64 aa. 40 (Dp) Related Amino Acid Sequences - Alignment No. 172 - gi No. 4758714 - Description: 45 - % Identity: 41.9 - Alignment Length: 131 - Location of Alignment in SEQ ID NO 264: from 1 to

50 Maximum Length Sequence: related to:

```
194
     Clone IDs:
           25093
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 265
5
          - Ceres seq id 1472772
          - Alternative transcription start site(s) located in SEQ
     ID NO 265:
            -1,3
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 266
            - Ceres seq id 1472773
            - Location of start within SEQ ID NO 265: at 3 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
15
     ID NO 266: at 32 aa.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Plant lipid transfer protein family
20
               - Location within SEQ ID NO 266: from 45 to 108 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 173
               - gi No. 3062791
25
               - Description:
               - % Identity:
                              72.2
               - Alignment Length: 90
               - Location of Alignment in SEO ID NO 266: from 21
     to 110
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 267
            - Ceres seq id 1472774
            - Location of start within SEQ ID NO 265: at 27 nt.
35
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 267: at 24 aa.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
40
                - Plant lipid transfer protein family
                - Location within SEQ ID NO 267: from 37 to 100 aa.
            (Dp) Related Amino Acid Sequences
                - Alignment No. 174
45
                - qi No. 3062791
                - Description:
                - % Identity: 72.2
```

- Alignment Length: 90

50

to 102

- Location of Alignment in SEQ ID NO 267: from 13

WO 00/40695 PCT/US00/00466 195 Maximum Length Sequence: related to: Clone IDs: 42300 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 268 - Ceres seq id 1533352 - Alternative transcription start site(s) located in SEQ ID NO 268: 10 17 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 269 - Ceres seq id 1533353 15 - Location of start within SEQ ID NO 268: at 1 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - haloacid dehalogenase-like hydrolase 20 - Location within SEQ ID NO 269: from 82 to 266 aa. (Dp) Related Amino Acid Sequences - Alignment No. 175 - qi No. 3913203 - Description: 25 - % Identity: 32.8 - Alignment Length: 196 - Location of Alignment in SEQ ID NO 269: from 81 to 266 30 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 270 - Ceres seq id 1533354 - Location of start within SEQ ID NO 268: at 25 nt. 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - haloacid dehalogenase-like hydrolase - Location within SEQ ID NO 270: from 74 to 258 aa. 40 (Dp) Related Amino Acid Sequences - Alignment No. 176 - gi No. 3913203 - Description: 45 - % Identity: 32.8

50 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 271

to 258

- Alignment Length: 196

- Location of Alignment in SEQ ID NO 270: from 73

196

- Ceres seq id 1533355
- Location of start within SEQ ID NO 268: at 265 nt.
- (C) Nomination and Annotation of Domains within
  5 Predicted Polypeptide(s)
  - haloacid dehalogenase-like hydrolase
  - Location within SEQ ID NO 271: from 1 to 178 aa.
  - (Dp) Related Amino Acid Sequences
- Alignment No. 177
  - qi No. 3913203
  - Description:
  - % Identity: 32.8
  - Alignment Length: 196
- Location of Alignment in SEQ ID NO 271: from 1 to 178

Maximum Length Sequence:

related to:

20 Clone IDs:

158412

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 272
  - Ceres seq id 1534544

25

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 273
  - Ceres seq id 1534545
- 30 Location of start within SEQ ID NO 272: at 2 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 178
        - qi No. 2317676
        - Description:
        - % Identity: 37.2
        - Alignment Length: 115
- Location of Alignment in SEQ ID NO 273: from 22 to 134
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 274
- 45 Ceres seg id 1534546
  - Location of start within SEQ ID NO 272: at 524 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 50 (Dp) Related Amino Acid Sequences
  - Alignment No. 179

197

- qi No. 4836939
- Description:
- % Identity: 47.3
- Alignment Length: 150
- 5 Location of Alignment in SEQ ID NO 274: from 19 to 166
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 275
- 10 Ceres seq id 1534547
  - Location of start within SEQ ID NO 272: at 815 nt.
  - (C) Nomination and Annotation of Domains within  $Predicted\ Polypeptide(s)$ 
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 180
      - gi No. 4836939
      - Description:
      - % Identity: 47.3
- 20 Alignment Length: 150
  - Location of Alignment in SEQ ID NO 275: from 1 to
- (Ba) Polypeptide Activities: Similar to calcium independent phrophospholipase A2 protein activities, and acy-protein thioesterase activities.

Maximum Length Sequence:

30 related to:

69

15

Clone IDs:

108109

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 276
- 35 Ceres seq id 1567172
  - Alternative transcription start site(s) located in SEQ ID NO 276:

- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 277
  - Ceres seq id 1567173
  - Location of start within SEQ ID NO 276: at 152 nt.
- 45 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 181
    - gi No. 1076301

198 - Description: CONSTANS protein - Arabidopsis thaliana >gi|1161514|emb|CAA64407| (X94937) CONSTANS protein [Arabidopsis thaliana] - % Identity: 49.2 - Alignment Length: 63 - Location of Alignment in SEQ ID NO 277: from 12 to 74 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 278 - Ceres seq id 1567174 - Location of start within SEQ ID NO 276: at 155 nt. (C) Nomination and Annotation of Domains within 15 Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 182 - qi No. 1076301 - Description: CONSTANS protein - Arabidopsis thaliana >gi|1161514|emb|CAA64407| (X94937) CONSTANS protein 20 [Arabidopsis thaliana] - % Identity: 49.2 - Alignment Length: 63 - Location of Alignment in SEQ ID NO 278: from 11 25 to 73 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 279 - Ceres seq id 1567175 30 - Location of start within SEQ ID NO 276: at 443 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 35 - Alignment No. 183 - qi No. 1076301 - Description: CONSTANS protein - Arabidopsis thaliana >qi|1161514|emb|CAA64407| (X94937) CONSTANS protein [Arabidopsis thaliana] 40 - % Identity: 67.4 - Alignment Length: 43 - Location of Alignment in SEQ ID NO 279: from 261 to 303 45 Maximum Length Sequence: related to:

Clone IDs: 168 Public Genomic DNA: 50 qi No: 4895213 Predicted Exons:

5

199

INIT 22484 ... 22368 OCKHAMG-CDS TERM 21760 ... 21704 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 280
- Ceres seq id 1567535
- Alternative transcription start site(s) located in SEQ ID NO 280:

12, 22, 23, 25, 28, 36, 68

- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 281
  - Ceres seq id 1567536
  - Location of start within SEQ ID NO 280: at 104 nt.
- 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 184
    - gi No. 1351365
- Description: UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 6.7 KD PROTEIN (CR6) >gi|2130002|pir||S68969 ubiquinol--cytochrome-c reductase (EC 1.10.2.2) potato >gi|633683|emb|CAA57768| (X82325) cytochrome c reductase subunit [Solanum tuberosum]
- 25 % Identity: 52.6
  - Alignment Length: 58
  - Location of Alignment in SEQ ID NO 281: from 2 to
- 30 Maximum Length Sequence:

related to:

Clone IDs:

57

21305

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 282
    - Ceres seq id 1569689
- Alternative transcription start site(s) located in SEQ ID NO 282:

28,29,49

40

35

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 283
  - Ceres seq id 1569690
  - Location of start within SEQ ID NO 282: at 1 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 283: at 32 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 50 (Dp) Related Amino Acid Sequences

```
200
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 284
            - Ceres seq id 1569691
            - Location of start within SEQ ID NO 282: at 92 nt.
 5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 185
               - qi No. 2191138
10
               - Description: (AF007269) A IG002N01.18 gene
     product [Arabidopsis thaliana]
               - % Identity: 19.5
               - Alignment Length: 87
15
               - Location of Alignment in SEQ ID NO 284: from 3 to
     83
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 285
20
            - Ceres seq id 1569692
            - Location of start within SEQ ID NO 282: at 248 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
25
           (Dp) Related Amino Acid Sequences
               - Alignment No. 186
               - qi No. 2191138
               - Description: (AF007269) A IG002N01.18 gene
     product [Arabidopsis thaliana]
               - % Identity:
30
               - Alignment Length: 87
               - Location of Alignment in SEQ ID NO 285: from 1 to
     31
35
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
     Maximum Length Sequence:
      related to:
40
     Clone IDs:
           41988
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 286
          - Ceres seq id 1571042
45
          - Alternative transcription start site(s) located in SEQ
     ID NO 286:
            -1,2,3,4,7
```

(B) Polypeptide Sequence

50

- Pat. Appln. SEQ ID NO 287

- Ceres seq id 1571043

201

- Location of start within SEQ ID NO 286: at 63 nt.

- (C) Nomination and Annotation of Domains within  $Predicted\ Polypeptide(s)$ 
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 187
    - qi No. 2129641
    - Description: major latex protein type 1 -

Arabidopsis thaliana >gi|1107493|emb|CAA63026| (X91960) major 10 latex protein type1 [Arabidopsis thaliana]

- % Identity: 71.3
- Alignment Length: 150
- Location of Alignment in SEQ ID NO 287: from 1 to

150

15

5

Maximum Length Sequence:

related to:

Clone IDs:

16143

- 20 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 288
  - Ceres seq id 1571079
  - Alternative transcription start site(s) located in SEQ ID NO 288:
- 25 -5,37,86,341,350
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 289
    - Ceres seq id 1571080
- 30 Location of start within SEQ ID NO 288: at 51 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Pathogenesis-related protein Bet v I family
- Location within SEQ ID NO 289: from 5 to 155 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 188
    - qi No. 1321731
- Description: (Z72439) major allergen Cor a 1

[Corylus avellana]

- % Identity: 35.7
- Alignment Length: 159
- Location of Alignment in SEQ ID NO 289: from 5 to

45 155

Maximum Length Sequence:

related to:

Clone IDs:

50 42101

(Ac) cDNA Polynucleotide Sequence

202

- Pat. Appln. SEQ ID NO 290
- Ceres seq id 1572097
- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 291
  - Ceres seq id 1572098
  - Location of start within SEQ ID NO 290: at 63 nt.
- 10 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Adhesion lipoprotein
  - Location within SEQ ID NO 291: from 118 to 191

aa.

15

- (Dp) Related Amino Acid Sequences
  - Alignment No. 189
  - gi No. 2129641
  - Description: major latex protein type 1 -
- Arabidopsis thaliana >gi|1107493|emb|CAA63026| (X91960) major latex protein typel [Arabidopsis thaliana]
  - % Identity: 69.5
  - Alignment Length: 154
  - Location of Alignment in SEQ ID NO 291: from 1 to
- 25 154
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 292
  - Ceres seq id 1572099
- Location of start within SEQ ID NO 290: at 336 nt.
  - - Adhesion lipoprotein
- Location within SEQ ID NO 292: from 27 to 100 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 190
    - qi No. 2129641
- 40 Description: major latex protein type 1 -

Arabidopsis thaliana >gi|1107493|emb|CAA63026| (X91960) major latex protein typel [Arabidopsis thaliana]

- % Identity: 69.5
- Alignment Length: 154
- 45 Location of Alignment in SEQ ID NO 292: from 1 to

63

Maximum Length Sequence:

related to:

50 Clone IDs:

203

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 293
- Ceres seq id 1572890
- Alternative transcription start site(s) located in SEQ 5 ID NO 293:

-10

15

20

35

40

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 294
- 10 Ceres seq id 1572891
  - Location of start within SEQ ID NO 293: at 1 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Photosystem I psaG / psaK
    - Location within SEQ ID NO 294: from 71 to 151 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 191
    - gi No. 3885511
  - Description: (AF084200) similar to PSI-K subunit of photosystem I from barley [Medicago sativa]
    - % Identity: 80.3
    - Alignment Length: 128
- 25 Location of Alignment in SEQ ID NO 294: from 22 to 148
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 295
- 30 Ceres seg id 1572892
  - Location of start within SEQ ID NO 293: at 64 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Photosystem I psaG / psaK
    - Location within SEO ID NO 295: from 50 to 130 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 192
      - qi No. 3885511
  - Description: (AF084200) similar to PSI-K subunit of photosystem I from barley [Medicago sativa]
    - % Identity: 80.3
    - Alignment Length: 128
- Location of Alignment in SEQ ID NO 295: from 1 to 127
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 296
    - Ceres seq id 1572893
    - Location of start within SEQ ID NO 293: at 76 nt.

204

(C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Photosystem I psaG / psaK 5 - Location within SEQ ID NO 296: from 46 to 126 aa. (Dp) Related Amino Acid Sequences - Alignment No. 193 - gi No. 3885511 - Description: (AF084200) similar to PSI-K subunit 10 of photosystem I from barley [Medicago sativa] - % Identity: 80.3 - Alignment Length: 128 - Location of Alignment in SEQ ID NO 296: from 1 to 15 123 Maximum Length Sequence: related to: Clone IDs: 20 33027 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 297 - Ceres seq id 1573606 - Alternative transcription start site(s) located in SEQ 25 ID NO 297: 7,8,13 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 298 30 - Ceres seq id 1573607 - Location of start within SEQ ID NO 297: at 56 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 298: at 19 aa. 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Pollen proteins Ole e I family - Location within SEQ ID NO 298: from 32 to 131 aa. 40 (Dp) Related Amino Acid Sequences - Alignment No. 194 - qi No. 4584110 - Description: (AJ133639) SAH7 protein [Arabidopsis thaliana] 45 - % Identity: 48.5 - Alignment Length: 163 - Location of Alignment in SEQ ID NO 298: from 1 to 163 50 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 299

```
205
            - Ceres seq id 1573608
            - Location of start within SEQ ID NO 297: at 71 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 299: at 14 aa.
 5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Pollen proteins Ole e I family
               - Location within SEQ ID NO 299: from 27 to 126 aa.
10
           (Dp) Related Amino Acid Sequences
               - Alignment No. 195
               - gi No. 4584110
               - Description: (AJ133639) SAH7 protein [Arabidopsis
15
     thaliana]
               - % Identity: 48.5
               - Alignment Length: 163
               - Location of Alignment in SEQ ID NO 299: from 1 to
     158
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 300
            - Ceres seq id 1573609
            - Location of start within SEQ ID NO 297: at 107 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Pollen proteins Ole e I family
               - Location within SEQ ID NO 300: from 15 to 114 aa.
30
           (Dp) Related Amino Acid Sequences
               - Alignment No. 196
               - qi No. 4584110
               - Description: (AJ133639) SAH7 protein [Arabidopsis
35
     thaliana]
               - % Identity: 48.5
               - Alignment Length: 163
               - Location of Alignment in SEQ ID NO 300: from 1 to
     146
40
    Maximum Length Sequence:
      related to:
     Clone IDs:
           31422
45
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 301
          - Ceres seq id 1573861
```

50 (B) Polypeptide Sequence
- Pat. Appln. SEQ ID NO 302

206

- Ceres seq id 1573862
- Location of start within SEQ ID NO 301: at 67 nt.
- (C) Nomination and Annotation of Domains within
  Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 197
    - qi No. 2191138
    - Description: (AF007269) A IG002N01.18 gene
- 10 product [Arabidopsis thaliana]
  - % Identity: 28.1
  - Alignment Length: 160
  - Location of Alignment in SEQ ID NO 302: from 3 to

156

15

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 303
  - Ceres seq id 1573863
  - Location of start within SEQ ID NO 301: at 223 nt.

20

25

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 198
    - gi No. 2191138
      - Description: (AF007269) A IG002N01.18 gene

product [Arabidopsis thaliana]

- % Identity: 28.1
- Alignment Length: 160
- Location of Alignment in SEQ ID NO 303: from 1 to
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

35

Maximum Length Sequence:

related to:

Clone IDs:

- 40 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 304
  - Ceres seq id 1574093
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 305
  - Ceres seq id 1574094
  - Location of start within SEQ ID NO 304: at 47 nt.
- 50 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

207

```
- Zinc finger, C3HC4 type (RING finger)
```

- Location within SEQ ID NO 305: from 152 to 192

aa.

```
5 (Dp) Related Amino Acid Sequences
```

- Alignment No. 199
- qi No. 3790593
- Description: (AF079185) RING-H2 finger protein

RHY1a [Arabidopsis thaliana]

- % Identity: 55.1
- Alignment Length: 49
- Location of Alignment in SEQ ID NO 305: from 148

to 196

10

- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 306
  - Ceres seq id 1574095
  - Location of start within SEQ ID NO 304: at 56 nt.
- 20 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Zinc finger, C3HC4 type (RING finger)
  - Location within SEQ ID NO 306: from 149 to 189

aa.

(Dp) Related Amino Acid Sequences

- Alignment No. 200
- qi No. 3790593
- Description: (AF079185) RING-H2 finger protein
- 30 RHYla [Arabidopsis thaliana]
  - % Identity: 55.1
  - Alignment Length: 49
  - Location of Alignment in SEQ ID NO 306: from 145

to 193

35

25

Maximum Length Sequence:

related to:

Clone IDs:

35447

40 Public Genomic DNA:

gi No: 4263694

Predicted Exons:

INIT 79640 ... 79806 GENBANK TERM 79888 ... 80164 GENBANK

45

INTR 79508 ... 79806 OCKHAMG-CDNA INTR 79888 ... 80309 OCKHAMG-CDNA

INIT 79640 ... 79806 OCKHAMG-CDS

50 TERM 79888 ... 80164 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

208

- Pat. Appln. SEQ ID NO 307
- Ceres seq id 1580388
- Alternative transcription start site(s) located in SEQ ID NO 307:

2, 3, 4, 5, 6, 7, 63, 94

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 308
  - Ceres seq id 1580389
  - Location of start within SEQ ID NO 307: at 133 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 308: at 27 aa.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Cystatin domain
  - Location within SEQ ID NO 308: from 87 to 141 aa.
  - (Dp) Related Amino Acid Sequences

- Alignment No. 201

- qi No. 2204077
- Description: (D85623) extracellular insoluble cystatin [Daucus carota]
  - % Identity: 44.9
  - Alignment Length: 136
- Location of Alignment in SEQ ID NO 308: from 6 to 141
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 309
    - Ceres seq id 1580390
    - Location of start within SEQ ID NO 307: at 142 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 309: at 24 aa.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Cystatin domain
  - Location within SEQ ID NO 309: from 84 to 138 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 202
    - qi No. 2204077
    - Description: (D85623) extracellular insoluble
- 45 cystatin [Daucus carota]
  - % Identity: 44.9
  - Alignment Length: 136
  - Location of Alignment in SEQ ID NO 309: from 3 to 138

50

Maximum Length Sequence:

PC1/US00/00460

```
209
      related to:
     Clone IDs:
           22677
     Public Genomic DNA:
 5
           gi No: 4972043
           Predicted Exons:
              SINGLE 18935 ... 19282
                                       GENBANK
              INTR 18856 ... 19527
                                       OCKHAMG-CDNA
10
              SINGLE 18935 ... 19282
                                       OCKHAMG-CDS
           gi No: 4972077
           Predicted Exons:
              INTR 60360 ... 61031 OCKHAMG-CDNA
15
              SINGLE 60439 ... 60786
                                       OCKHAMG-CDS
           gi No: 5748495
           Predicted Exons:
              INTR 60360 ... 61031 OCKHAMG-CDNA
20
              SINGLE 60439 ... 60786
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 310
          - Ceres seq id 1582959
25
          - Alternative transcription start site(s) located in SEO
     ID NO 310:
           2,5,6,7,58,61
        (B) Polypeptide Sequence
30
            - Pat. Appln. SEQ ID NO 311
            - Ceres seq id 1582960
            - Location of start within SEQ ID NO 310: at 80 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 311: at 19 aa.
35
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Dehydrins
               - Location within SEQ ID NO 311: from 37 to 113 aa.
40
           (Dp) Related Amino Acid Sequences
               - Alignment No. 203
               - qi No. 4972049
               - Description: (AL078470) glycine-rich protein like
45
     [Arabidopsis thaliana]
               - % Identity:
                              100
               - Alignment Length: 115
              - Location of Alignment in SEQ ID NO 311: from 1 to
     115
50
```

Maximum Length Sequence:

```
210
     related to:
    Clone IDs:
          15190
          34118
 5
    Public Genomic DNA:
          qi No: 4582444
          Predicted Exons:
                   39128 ... 39157 GENBANK
             TINI
             INTR 39241 ... 39388 GENBANK
10
             INTR 39469 ... 39640 GENBANK
             INTR 39922 ... 40108 GENBANK
             TERM 40198 ... 40413 GENBANK
             INTR 39945 ... 40108 OCKHAMG-CDNA
15
             INTR 40198 ... 40660 OCKHAMG-CDNA
             INTR 39054 ... 39157 OCKHAMG-CDNA
             INTR 39241 ... 39388 OCKHAMG-CDNA
             INTR 39469 ... 39640 OCKHAMG-CDNA
             INTR 39922 ... 40108 OCKHAMG-CDNA
20
             INTR 40198 ... 40626 OCKHAMG-CDNA
                   39128 ... 39157 OCKHAMG-CDS
             INIT
             INTR 39241 ... 39388 OCKHAMG-CDS
25
             INTR 39469 ... 39640 OCKHAMG-CDS
                   39922 ... 40108 OCKHAMG-CDS
             INTR
             TERM 40198 .. 40413 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
         - Pat. Appln. SEQ ID NO 312
30
         - Ceres seq id 1663221
         - Alternative transcription start site(s) located in SEO
    ID NO 312:
           -5, 2, 3, 4, 5, 12, 40, 448
         - Clone 15190 starts at 448 and ends at 1076 in cDNA.
35
         - Clone 34118 starts at 1 and ends at 1042 in cDNA.
       (B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 313
           - Ceres seq id 1663222
40
           - Location of start within SEQ ID NO 312: at 75 nt.
          (C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
              - Ribosomal protein S3, C-terminal domain.
45
              - Location within SEQ ID NO 313: from 104 to 188
    aa.
          (Dp) Related Amino Acid Sequences
```

- Alignment No. 204
- 50 gi No. 133940

211

- Description: 40S RIBOSOMAL PROTEIN S3A (S1A) >gi|70851|pir||R3XL3A ribosomal protein S3a - African clawed frog >gi|65091|emb|CAA40592| (X57322) ribosomal protein S1a [Xenopus laevis]

- % Identity: 80.1
- Alignment Length: 231
- Location of Alignment in SEQ ID NO 313: from 1 to 228
- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 314
  - Ceres seq id 1663223
  - Location of start within SEQ ID NO 312: at 204 nt.
- 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 314: from 61 to 145 aa.
- 20 (Dp) Related Amino Acid Sequences
  - Alignment No. 205
  - qi No. 133940
  - Description: 40S RIBOSOMAL PROTEIN S3A (S1A)

>gi|70851|pir||R3XL3A ribosomal protein S3a - African clawed frog >gi|65091|emb|CAA40592| (X57322) ribosomal protein S1a [Xenopus laevis]

- % Identity: 80.1
- Alignment Length: 231
- Location of Alignment in SEQ ID NO 314: from 1 to

30 185

25

35

5

Maximum Length Sequence:

related to:

Clone IDs:

3996

2749

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 315
  - Ceres seq\_id 1663275
- Alternative transcription start site(s) located in SEQ ID NO 315:
  - 3,4,5,6,7,8,9,15
  - Clone 3996 starts at 1 and ends at 484 in cDNA.
  - Clone 2749 starts at 7 and ends at 488 in cDNA.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 316
    - Ceres seq id 1663276
    - Location of start within SEQ ID NO 315: at 75 nt.

50

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212
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein S14p/S29e
               - Location within SEQ ID NO 316: from 3 to 54 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 206
               - qi No. 4506717
               - Description: ref | NP 001023.1 | pRPS29 | ribosomal
     protein S29 >qi|266972|sp|P30054|RS29 HUMAN 40S RIBOSOMAL
     PROTEIN S29 >qi|631884|pir||S30298 ribosomal protein S29 -
     rat >gi|1362934|pir||S55919 ribosomal protein S29 - human
     protein S29 [Homo sapiens]
               - % Identity: 72.2
               - Alignment Length: 54
               - Location of Alignment in SEQ ID NO 316: from 1 to
     54
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 317
            - Ceres seq id 1663277
            - Location of start within SEQ ID NO 315: at 92 nt.
           (C) Nomination and Annotation of Domains within
25
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
     Maximum Length Sequence:
      related to:
30
     Clone IDs:
           114940
           25068
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NC 318
          - Ceres seq id 1665304
          - Alternative transcription start site(s) located in SEQ
     ID NO 318:
            -8, -3, -1, 2, 3, 4, 8, 17, 251
          - Clone 114940 starts at 1 and ends at 743 in cDNA.
          - Clone 25068 starts at 4 and ends at 760 in cDNA.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 319
            - Ceres seq id 1665305
            - Location of start within SEQ ID NO 318: at 63 nt.
```

- 45
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 207
      - qi No. 2129641

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213 - Description: major latex protein type 1 -Arabidopsis thaliana >gi|1107493|emb|CAA63026| (X91960) major latex protein type1 [Arabidopsis thaliana] - % Identity: 71.3 5 - Alignment Length: 150 - Location of Alignment in SEQ ID NO 319: from 1 to 150 Maximum Length Sequence: 10 related to: Clone IDs: 94673 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 320 15 - Ceres seq id 1709970 - Alternative transcription start site(s) located in SEO ID NO 320: -2, -1, 11, 41, 42, 43, 44, 45, 46, 47, 48, 5120 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 321 - Ceres seq id 1709971 - Location of start within SEQ ID NO 320: at 276 nt. 25 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 208 - qi No. 2961300 30 - Description: (AJ225027) ribosomal protein L24 [Cicer arietinum] - % Identity: 86.2 - Alignment Length: 160 - Location of Alignment in SEQ ID NO 321: from 1 to 35 107 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 322 - Ceres seq id 1709972 40 - Location of start within SEQ ID NO 320: at 424 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 45 Maximum Length Sequence: related to: Clone IDs: 112835 (Ac) cDNA Polynucleotide Sequence 50

- Pat. Appln. SEQ ID NO 323

```
214
          - Ceres seq id 1711273
          - Alternative transcription start site(s) located in SEQ
     ID NO 323:
            2,28,30,33,56,62,235
 5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 324
            - Ceres seq id 1711274
            - Location of start within SEQ ID NO 323: at 92 nt.
10
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Helix-turn-helix
               - Location within SEQ ID NO 324: from 87 to 141 aa.
15
           (Dp) Related Amino Acid Sequences
               - Alignment No. 209
               - qi No. 1632831
               - Description: (Z49698) orf [Ricinus communis]
               - % Identity: 81
20
               - Alignment Length: 142
               - Location of Alignment in SEQ ID NO 324: from 1 to
     142
25
     Maximum Length Sequence:
      related to:
     Clone IDs:
           17878
     (Ac) cDNA Polynucleotide Sequence
30
          - Pat. Appln. SEQ ID NO 325
          - Ceres seq id 1715423
          - Alternative transcription start site(s) located in SEQ
     ID NO 325:
            2,4,7,66
35
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 326
            - Ceres seq id 1715424
            - Location of start within SEQ ID NO 325: at 25 nt.
40
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 210
45
               - qi No. 4337175
               - Description: (AC006416) ESTs gb|T20589,
     gb|T04648, gb|AA597906, gb|T04111, gb|R84180, gb|R65428,
     qb|T44439, qb|T76570, qb|R90004, qb|T45020, qb|T42457,
     gb|T20921, gb|AA042762 and gb|AA720210 come from this gene.
```

50

[Arabidopsis thaliana]

- % Identity: 44.8

215

- Alignment Length: 67

- Location of Alignment in SEQ ID NO 326: from 15 to 81

- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 327
  - Ceres seq id 1715425
  - Location of start within SEQ ID NO 325: at 40 nt.
- 10 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 211
    - qi No. 4337175
- 15 - Description: (AC006416) ESTs gb|T20589, gb|T04648, gb|AA597906, gb|T04111, gb|R84180, gb|R65428, gb|T44439, gb|T76570, gb|R90004, gb|T45020, gb|T42457, gb|T20921, gb|AA042762 and gb|AA720210 come from this gene. [Arabidopsis thaliana] 20
  - % Identity: 44.8
    - Alignment Length: 67
  - Location of Alignment in SEQ ID NO 327: from 10 to 76
- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 328
  - Ceres seq id 1715426
  - Location of start within SEQ ID NO 325: at 260 nt.
- 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 212
    - gi No. 4337175
- 35 - Description: (AC006416) ESTs qb/T20589, gb|T04648, gb|AA597906, gb|T04111, gb|R84180, gb|R65428, gb|T44439, gb|T76570, gb|R90004, gb|T45020, gb|T42457, gb|T20921, gb|AA042762 and gb|AA720210 come from this gene. [Arabidopsis thaliana]
  - % Identity: 56.1
  - Alignment Length: 57
  - Location of Alignment in SEQ ID NO 328: from 5 to

61

40

45 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene.

Maximum Length Sequence: related to:

50 Clone IDs:

216

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 329
- Ceres seq id 1715962
- Alternative transcription start site(s) located in SEQ 5 ID NO 329:

2,7,9,10,26,28,30,31,32,38,39,42,43,47,48,194,247

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 330
- Ceres seq id 1715963
  - Location of start within SEQ ID NO 329: at 87 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Ribosomal protein L6
    - Location within SEQ ID NO 330: from 12 to 194 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 213
    - gi No. 1710581
  - Description: 60S RIBOSOMAL PROTEIN L9 >gi|2129720|pir||S71255 ribosomal protein L9 Arabidopsis thaliana >gi|1107489|emb|CAA63024| (X91958) 60S ribosomal protein L9 [Arabidopsis thaliana]
    - % Identity: 93.3
    - Alignment Length: 195
  - Location of Alignment in SEQ ID NO 330: from 1 to 194
- 30 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 331
  - Ceres seq id 1715964
  - Location of start within SEQ ID NO 329: at 114 nt.
- 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein L6
  - Location within SEQ ID NO 331: from 3 to 185 aa.
- 40 (Dp) Related Amino Acid Sequences
  - Alignment No. 214
  - gi No. 1710581
  - Description: 60S RIBOSOMAL PROTEIN L9
- >gi|2129720|pir||S71255 ribosomal protein L9 Arabidopsis thaliana >gi|1107489|emb|CAA63024| (X91958) 60S ribosomal protein L9 [Arabidopsis thaliana]
  - % Identity: 93.3
  - Alignment Length: 195
  - Location of Alignment in SEQ ID NO 331: from 1 to
- 50 185

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217

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(B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 332
           - Ceres seq id 1715965
           - Location of start within SEQ ID NO 329: at 369 nt.
5
           (C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
              - Ribosomal protein L6
              - Location within SEQ ID NO 332: from 1 to 100 aa.
10
           (Dp) Related Amino Acid Sequences
              - Alignment No. 215
              - qi No. 1710581
              - Description: 60S RIBOSOMAL PROTEIN L9
    >gi|2129720|pir||S71255 ribosomal protein L9 - Arabidopsis
15
    thaliana >gi|1107489|emb|CAA63024| (X91958) 60S ribosomal
    protein L9 [Arabidopsis thaliana]
              - % Identity:
                             93.3
              - Alignment Length: 195
20
              - Location of Alignment in SEQ ID NO 332: from 1 to
    100
    Maximum Length Sequence:
     related to:
25
    Clone IDs:
          41712
    Public Genomic DNA:
          qi No: 4512656
          Predicted Exons:
             INIT 106774... 106633 GENBANK
30
             INTR 106277... 106194 GENBANK
             INTR 105920... 105838 GENBANK
             TERM 105749... 105630 GENBANK
35
             INTR 106841... 106633 OCKHAMG-CDNA
             INTR 106277... 106194 OCKHAMG-CDNA
             INTR 105920... 105838 OCKHAMG-CDNA
             INTR 105749... 105453 OCKHAMG-CDNA
40
             INIT 106774... 106633 OCKHAMG-CDS
             INTR 106277... 106194 OCKHAMG-CDS
             INTR 105920... 105838 OCKHAMG-CDS
                    105749... 105630 OCKHAMG-CDS
             TERM
           qi No: 4559319
45
           Predicted Exons:
              INIT 10855 ... 10714
                                       GENBANK
              INTR 10358 ... 10275
                                       GENBANK
             INTR 10001 ... 9919
TERM 9830 ... 9711
                                       GENBANK
                                       GENBANK
50
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INTR 10922 ... 10714 OCKHAMG-CDNA

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218
              INTR 10358 ...
                               10275 OCKHAMG-CDNA
                               9919
9534
              INTR
                    10001 ...
                                        OCKHAMG-CDNA
              INTR 9830 ...
                                        OCKHAMG-CDNA
5
                    10855 ... 10714 OCKHAMG-CDS
              INIT
              INTR
                    10358 ... 10275 OCKHAMG-CDS
              INTR
                    10001 ... 9919
                                        OCKHAMG-CDS
                   9830 ... 9711
              TERM
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
10
          - Pat. Appln. SEQ ID NO 333
          - Ceres seq id 1808584
          - Alternative transcription start site(s) located in SEQ
    ID NO 333:
            34
15
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 334
            - Ceres seq id 1808585
            - Location of start within SEQ ID NO 333: at 1 nt.
20
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Helix-turn-helix
               - Location within SEQ ID NO 334: from 110 to 164
25
     aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 216
               - qi No. 1632831
30
               - Description: (Z49698) orf [Ricinus communis]
               - % Identity: 81
               - Alignment Length: 142
               - Location of Alignment in SEQ ID NO 334: from 24
     to 165
35
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 335
            - Ceres seq id 1808586
            - Location of start within SEQ ID NO 333: at 70 nt.
40
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Helix-turn-helix
               - Location within SEQ ID NO 335: from 87 to 141 aa.
45
           (Dp) Related Amino Acid Sequences
               - Alignment No. 217
               - qi No. 1632831
               - Description: (Z49698) orf [Ricinus communis]
50
               - % Identity: 81
               - Alignment Length: 142
```

219

- Location of Alignment in SEQ ID NO 335: from 1 to

142

Maximum Length Sequence:

5 related to:

Clone IDs:

42211

Public Genomic DNA:

gi No: 4914400

10 Predicted Exons:

INTR 90159 ... 89532 OCKHAMG-CDNA

SINGLE 90124 ... 89615 OCKHAMG-CDS

gi No: 4938473

15 Predicted Exons:

SINGLE 2368 ... 1859 GENBANK

INTR 2403 ... 1776 OCKHAMG-CDNA

20 SINGLE 2368 ... 1859 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 336
- Ceres seq id 1808591
- Alternative transcription start site(s) located in SEQ

25 ID NO 336:

7,10

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 337
- 30 Ceres seq id 1808592
  - Location of start within SEQ ID NO 336: at 2 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 337: at 36 aa.
- 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 218
    - qi No. 4926823
- Description: (AC004135) T17H7.8 [Arabidopsis

thaliana]

- % Identity: 46.1
- Alignment Length: 178
- Location of Alignment in SEQ ID NO 337: from 13
- 45 to 177
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 338
    - Ceres seq id 1808593
- Location of start within SEQ ID NO 336: at 38 nt.

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220
            - Location of Signal Peptide Cleavage Site within SEO
    ID NO 338: at 24 aa.
           (C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 219
               - gi No. 4926823
               - Description: (AC004135) T17H7.8 [Arabidopsis
    thaliana |
               - % Identity: 46.1
               - Alignment Length: 178
               - Location of Alignment in SEQ ID NO 338: from 1 to
     165
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 339
            - Ceres seq id 1808594
            - Location of start within SEQ ID NO 336: at 164 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 220
               - gi No. 4926823
               - Description: (AC004135) T17H7.8 [Arabidopsis
     thaliana]
               - % Identity: 46.1
               - Alignment Length: 178
               - Location of Alignment in SEQ ID NO 339: from 1 to
     123
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot specific gene, plant specific gene.
35
     Maximum Length Sequence:
      related to:
     Clone IDs:
           2217
     (Ac) cDNA Polynucleotide Sequence
40
          - Pat. Appln. SEQ ID NO 340
          - Ceres seq id 1920563
          - Alternative transcription start site(s) located in SEO
     ID NO 340:
45
            6
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 341
```

- Location of start within SEQ ID NO 340: at 1 nt.

- Ceres seg id 1920564

221

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Oxidoreductase FAD/NAD-binding domain
  - Location within SEQ ID NO 341: from 45 to 167 aa.

5 (Dp) Related Amino Acid Sequences

- Alignment No. 221
- qi No. 729477
- Description: FERREDOXIN--NADP REDUCTASE PRECURSOR (FNR) >gi|320548|pir||A44974 ferredoxin--NADP+ reductase (EC 1.18.1.2) precursor common ice plant >gi|167256 (M25528) ferredoxin-NADP+ reductase precursor (fnrA; EC 1.6.7.1) [Mesembryanthemum
  - % Identity: 89.8
  - Alignment Length: 197
  - Location of Alignment in SEQ ID NO 341: from 1 to 197
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 342
      - Ceres seq id 1920565
      - Location of start within SEQ ID NO 340: at 112 nt.
- (C) Nomination and Annotation of Domains within 25 Predicted Polypeptide(s)
  - Oxidoreductase FAD/NAD-binding domain
  - Location within SEQ ID NO 342: from 8 to 130 aa.
  - (Dp) Related Amino Acid Sequences
  - Alignment No. 222
    - qi No. 729477
  - Description: FERREDOXIN--NADP REDUCTASE PRECURSOR (FNR) >gi|320548|pir||A44974 ferredoxin--NADP+ reductase (EC 1.18.1.2) precursor common ice plant >gi|167256 (M25528)
- ferredoxin-NADP+ reductase precursor (fnrA; EC 1.6.7.1)
  [Mesembryanthemum
  - % Identity: 89.8
  - Alignment Length: 197
  - Location of Alignment in SEQ ID NO 342: from 1 to
- 40 160

15

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 343
  - Ceres seq id 1920566
- 45 Location of start within SEQ ID NO 340: at 118 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
    - Oxidoreductase FAD/NAD-binding domain
- Location within SEQ ID NO 343: from 6 to 128 aa.

222

```
(Dp) Related Amino Acid Sequences
               - Alignment No. 223
               - qi No. 729477
               - Description: FERREDOXIN--NADP REDUCTASE PRECURSOR
5
     (FNR) >gi|320548|pir||A44974 ferredoxin--NADP+ reductase (EC
    1.18.1.2) precursor - common ice plant >qi|167256 (M25528)
    ferredoxin-NADP+ reductase precursor (fnrA; EC 1.6.7.1)
     [Mesembryanthemum
              - % Identity: 89.8
10
               - Alignment Length: 197
               - Location of Alignment in SEQ ID NO 343: from 1 to
    158
    Maximum Length Sequence:
15
     related to:
    Clone IDs:
          13864
    Public Genomic DNA:
          qi No: 4309719
20
          Predicted Exons:
              INTT
                    50987 ... 50773 GENBANK
              INTR 49622 ... 49352
                                       GENBANK
              INTR 49257 ... 49162 GENBANK
              INTR 48896 ... 48786 GENBANK
25
              INTR 48686 ... 48377
                                      GENBANK
              INTR 48133 ... 48043 GENBANK
              INTR 47924 ... 47591 GENBANK
              INTR 47475 ...
                               47327
                                     GENBANK
              TERM 47084 ...
                               46550
                                       GENBANK
30
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 344
          - Ceres seq id 1974419
          - Alternative transcription start site(s) located in SEO
     ID NO 344:
35
            -5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 345
            - Ceres seq id 1974420
40
            - Location of start within SEQ ID NO 344: at 81 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - ATPases associated with various cellular
45
     activities (AAA)
               - Location within SEQ ID NO 345: from 454 to 664
     aa.
           (Dp) Related Amino Acid Sequences
```

- Alignment No. 224 - gi No. 4006905

223

- Description: (Z99708) ATPase-like protein [Arabidopsis thaliana]
  - % Identity: 84.3
  - Alignment Length: 633
- Location of Alignment in SEQ ID NO 345: from 72 to 700
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 346
- Ceres seq\_id 1974421
  - Location of start within SEQ ID NO 344: at 96 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- ATPases associated with various cellular activities (AAA)
  - Location within SEQ ID NO 346: from 449 to 659 aa.
- 20 (Dp) Related Amino Acid Sequences
  - Alignment No. 225
  - qi No. 4006905
  - Description: (Z99708) ATPase-like protein

[Arabidopsis thaliana]

- 25 % Identity: 84.3
  - Alignment Length: 633
  - Location of Alignment in SEQ ID NO 346: from 67 to 695
- 30 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 347
  - Ceres seq id 1974422
  - Location of start within SEQ ID NO 344: at 141 nt.
- 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - ATPases associated with various cellular activities (AAA)  $\,$
  - Location within SEQ ID NO 347: from 434 to 644
- 40 aa.

5

10

- (Dp) Related Amino Acid Sequences
  - Alignment No. 226
  - gi No. 4006905
- Description: (Z99708) ATPase-like protein

[Arabidopsis thaliana]

- % Identity: 84.3
- Alignment Length: 633
- Location of Alignment in SEQ ID NO 347: from 52
- 50 to 680

```
224
     Maximum Length Sequence:
      related to:
     Clone IDs:
           36333
 5
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 348
          - Ceres seq id 1975983
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 349
            - Ceres seq id 1975984
            - Location of start within SEQ ID NO 348: at 90 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 227
               - qi No. 3688432
20
               - Description: (AJ011705) 40S ribosomal protein S10
     [Lumbricus rubellus]
               - % Identity: 55.3
               - Alignment Length: 164
               - Location of Alignment in SEQ ID NO 349: from 32
25
     to 192
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 350
            - Ceres seq id 1975985
            - Location of start within SEQ ID NO 348: at 135 nt.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
35
               - Alignment No. 228
               - qi No. 3688432
               - Description: (AJ011705) 40S ribosomal protein S10
     [Lumbricus rubellus]
               - % Identity: 55.3
               - Alignment Length: 164
40
               - Location of Alignment in SEQ ID NO 350: from 17
     to 177
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 351
45
            - Ceres seq id 1975986
```

- Location of start within SEQ ID NO 348: at 183 nt.

(C) Nomination and Annotation of Domains within

(Dp) Related Amino Acid Sequences

50

Predicted Polypeptide(s)

225

- Alignment No. 229
- qi No. 3688432
- Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]
  - % Identity: 55.3
  - Alignment Length: 164
- Location of Alignment in SEQ ID NO 351: from 1 to 161
- (Ba) Polypeptide Activities: Similar to glycine-rich RNA binding ABA inducible protein activities.

Maximum Length Sequence:

related to:

Clone IDs:

CIONE IDS.

5

15

24587

41072

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 352
- Ceres seq id 1976019
- Alternative transcription start site(s) located in SEQ ID NO 352:

13

- Clone 24587 starts at 13 and ends at in cDNA.
- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 353
  - Ceres seq id 1976020
  - Location of start within SEQ ID NO 352: at 101 nt.
- 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 230
    - qi No. 3688432
  - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]
    - % Identity: 55.3
    - Alignment Length: 164
    - Location of Alignment in SEQ ID NO 353: from 1 to
- 40 161

35

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 354
  - Ceres seq id 1976021
- 45 Location of start within SEQ ID NO 352: at 239 nt.
  - - (Dp) Related Amino Acid Sequences
      - Alignment No. 231
      - gi No. 3688432

226

- Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]

- % Identity: 55.3
- Alignment Length: 164
- 5 Location of Alignment in SEQ ID NO 354: from 1 to 115
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 355
    - Ceres seq id 1976022
      - Location of start within SEQ ID NO 352: at 287 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 232
      - qi No. 3688432
  - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]
    - % Identity: 55.3
      - Alignment Length: 164
      - Location of Alignment in SEQ ID NO 355: from 1 to
- 25 (Ba) Polypeptide Activities: Extensin like protein activities and glycine rich protein activities.

Maximum Length Sequence:

related to:

30 Clone IDs:

99

10

15

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117263

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 356
  - Ceres seq id 1976673
- Alternative transcription start site(s) located in SEQ ID NO 356:
  - -46,2,3,6,7
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 357
    - Ceres seq id 1976674
    - Location of start within SEQ ID NO 356: at 3 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 357: at 34 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Ribosomal protein S21
    - Location within SEQ ID NO 357: from 106 to 155
- 50 aa.

40

227

- (Dp) Related Amino Acid Sequences
  - Alignment No. 233
  - qi No. 1710750
  - Description: 30S RIBOSOMAL PROTEIN S21 (BS-B)
- 5 >gi|1303814|dbj|BAA12470| (D84432) YqeX [Bacillus subtilis] >gi|1890063|dbj|BAA12082| (D83717) YqeX [Bacillus subtilis] >gi|2634987|emb|CAB14483| (Z99117) ribosomal protein S21 [Bacillus subtilis]
  - % Identity: 38.5
    - Alignment Length: 52
  - Location of Alignment in SEQ ID NO 357: from 106 to 157  $\,$ 
    - (B) Polypeptide Sequence

10

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- Pat. Appln. SEQ ID NO 358
- Ceres seq id 1976675
- Location of start within SEQ ID NO 356: at 48 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 358: at 19 aa.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S21
  - Location within SEQ ID NO 358: from 91 to 140 aa.
- 25 (Dp) Related Amino Acid Sequences
  - Alignment No. 234
  - qi No. 1710750
  - Description: 30S RIBOSOMAL PROTEIN S21 (BS-B)
- 30 >gi|1303814|dbj|BAA12470| (D84432) YqeX [Bacillus subtilis] >gi|1890063|dbj|BAA12082| (D83717) YqeX [Bacillus subtilis] >gi|2634987|emb|CAB14483| (Z99117) ribosomal protein S21 [Bacillus subtilis]
  - % Identity: 38.5
  - Alignment Length: 52
  - Location of Alignment in SEQ ID NO 358: from 91 to 142
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 359
      - Ceres seq id 1976676
      - Location of start within SEQ ID NO 356: at 192 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S21
  - Location within SEQ ID NO 359: from 43 to 92 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 235
    - qi No. 1710750

228

```
- Description: 30S RIBOSOMAL PROTEIN S21 (BS-B)
    >gi|1303814|dbj|BAA12470| (D84432) YqeX [Bacillus subtilis]
    >gi|1890063|dbj|BAA12082| (D83717) YqeX [Bacillus subtilis]
    >gi|2634987|emb|CAB14483| (Z99117) ribosomal protein S21
5
    [Bacillus subtilis]
              - % Identity: 38.5
              - Alignment Length: 52
              - Location of Alignment in SEQ ID NO 359: from 43
    to 94
10
    Maximum Length Sequence:
     related to:
    Clone IDs:
          42333
15
    Public Genomic DNA:
          qi No: 6041810
          Predicted Exons:
             INTR 93582 ... 94052 OCKHAMG-CDNA
             INTR 94169 ... 94771
                                     OCKHAMG-CDNA
20
             INIT 93630 ... 94052 OCKHAMG-CDS
             TERM 94169 ... 94687 OCKHAMG-CDS
          gi No: 6091711
          Predicted Exons:
25
             INTR 60479 ... 60949 OCKHAMG-CDNA
             INTR 61066 ... 61668 OCKHAMG-CDNA
             INIT 60527 ... 60949 OCKHAMG-CDS
             TERM 61066 ... 61584 OCKHAMG-CDS
30
          qi No: 6102641
          Predicted Exons:
             INTR 52165 ... 52635 OCKHAMG-CDNA
             INTR 52752 ... 53354
                                     OCKHAMG-CDNA
35
             INIT 52213 ... 52635 OCKHAMG-CDS
             TERM 52752 ... 53270 OCKHAMG-CDS
          gi No: 6453849
          Predicted Exons:
                   59432 ... 58964
             INTR
                                     OCKHAMG-CDNA
40
             INTR 58847 ... 58243
                                      OCKHAMG-CDNA
             INIT 59386 ... 58964 OCKHAMG-CDS
             TERM 58847 ...
                              58329
                                      OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
45
         - Pat. Appln. SEQ ID NO 360
         - Ceres seq id 2025186
        (B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 361

- Ceres seq id 2025187

229

- Location of start within SEQ ID NO 360: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - D-isomer specific 2-hydroxyacid dehydrogenases
  - Location within SEQ ID NO 361: from 93 to 295 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 236
    - gi No. 5458864
- Description: (AJ248287) GLYCERATE DEHYDROGENASE [Pyrococcus abyssi]
  - % Identity: 42.8
  - Alignment Length: 299
- Location of Alignment in SEQ ID NO 361: from 36 to 322
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 362
- 20 Ceres seq id 2025188

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- Location of start within SEQ ID NO 360: at 49 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - D-isomer specific 2-hydroxyacid dehydrogenases
  - Location within SEQ ID NO 362: from 77 to 279 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 237
    - qi No. 5458864
- Description: (AJ248287) GLYCERATE DEHYDROGENASE [Pyrococcus abyssi]
  - % Identity: 42.8
  - Alignment Length: 299
- Location of Alignment in SEQ ID NO 362: from 20 to 306
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 363
    - Ceres seq id 2025189
    - Location of start within SEQ ID NO 360: at 70 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - D-isomer specific 2-hydroxyacid dehydrogenases
  - Location within SEQ ID NO 363: from 70 to 272 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 238
- gi No. 5458864

230 - Description: (AJ248287) GLYCERATE DEHYDROGENASE [Pyrococcus abyssi] - % Identity: 42.8 - Alignment Length: 299 5 - Location of Alignment in SEQ ID NO 363: from 13 to 299 Maximum Length Sequence: related to: Clone IDs: 10 267004 Public Genomic DNA: qi No: 2924730 Predicted Exons: INTR 8331 ... 8882 OCKHAMG-CDNA 15 SINGLE 8366 ... 8686 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 364 - Ceres seq id 2025372 20 - Alternative transcription start site(s) located in SEQ ID NO 364: 21 (B) Polypeptide Sequence 25 - Pat. Appln. SEQ ID NO 365 - Ceres seg id 2025373 - Location of start within SEQ ID NO 364: at 1 nt. 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 239 - gi No. 132825 - Description: 50S RIBOSOMAL PROTEIN CL25, 35 CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein PsCL25 precursor, chloroplast - garden pea >qi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein (AA -30 to 74) [Pisum sativum] 40 - % Identity: 56.1 - Alignment Length: 82 - Location of Alignment in SEQ ID NO 365: from 14 to 95 (B) Polypeptide Sequence 45 - Pat. Appln. SEQ ID NO 366 - Ceres seq id 2025374 - Location of start within SEQ ID NO 364: at 37 nt.

- Location of Signal Peptide Cleavage Site within SEQ

50

ID NO 366: at 16 aa.

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(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 240
 5
               - qi No. 132825
               - Description: 50S RIBOSOMAL PROTEIN CL25,
     CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein
     PsCL25 precursor, chloroplast - garden pea
     >qi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein
10
     (AA - 30 \text{ to } 74) [Pisum sativum]
               - % Identity: 56.1
               - Alignment Length: 82
               - Location of Alignment in SEQ ID NO 366: from 2 to
     83
15
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 367
            - Ceres seq id 2025375
            - Location of start within SEQ ID NO 364: at 142 nt.
20
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 241
25
               - qi No. 132825
               - Description: 50S RIBOSOMAL PROTEIN CL25,
     CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein
     PsCL25 precursor, chloroplast - garden pea
     >qi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein
     (AA - 30 \text{ to } 74) [Pisum sativum]
30
               - % Identity: 56.1
               - Alignment Length: 82
               - Location of Alignment in SEQ ID NO 367: from 1 to
     48
35
     Maximum Length Sequence:
      related to:
     Clone IDs:
           26930
40
     Public Genomic DNA:
           qi No: 5225383
           Predicted Exons:
              SINGLE 64539 ... 64276 GENBANK
45
                     64787 ... 64692 OCKHAMG-CDS
              INIT
              TERM 64620 ... 64276 OCKHAMG-CDS
           gi No: 6136349
           Predicted Exons:
              INIT 126797... 126892 OCKHAMG-CDS
50
              TERM 126964... 127308 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
```

232

- Pat. Appln. SEQ ID NO 368
- Ceres seq id 2025471
- Alternative transcription start site(s) located in SEQ ID NO 368:
  - 95, 96, 97, 98, 100, 101, 102, 105, 106, 110
  - Clone 26930 starts at 97 and ends at 631 in cDNA.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 369
- Ceres seq id 2025472 10
  - Location of start within SEQ ID NO 368: at 1 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 242
      - qi No. 5732082
      - Description: (AF162444) contains similarity to

Solanum lycopersicum (tomato) wound induced protein

- (GB:X59882) [Arabidopsis thaliana] 20
  - % Identity: 100
  - Alignment Length: 87
  - Location of Alignment in SEQ ID NO 369: from 60

to 146

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 370
  - Ceres seq id 20254.3
  - Location of start within SEQ ID NO 368: at 178 nt.

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 243
- qi No. 5732082
  - Description: (AF162444) contains similarity to

Solanum lycopersicum (tomato) wound induced protein

(GB:X59882) [Arabidopsis thaliana]

- % Identity: 100
- Alignment Length: 87
- Location of Alignment in SEQ ID NO 370: from 1 to

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 371
  - Ceres seq id 2025474
  - Location of start within SEQ ID NO 368: at 214 nt.
- (C) Nomination and Annotation of Domains within 50 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences

PC1/US00/0046

233

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- Alignment No. 244
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- qi No. 5732082
- Description: (AF162444) contains similarity to

Solanum lycopersicum (tomato) wound induced protein

5 (GB:X59882) [Arabidopsis thaliana]

- % Identity: 100
- Alignment Length: 87
- Location of Alignment in SEQ ID NO 371: from 1 to

75

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Maximum Length Sequence:

related to:

Clone IDs:

27681

15 Public Genomic DNA:

gi No: 6434227

Predicted Exons:

INTR 51529 ... 51786 OCKHAMG-CDNA

INIT 47952 ... 48020 OCKHAMG-CDS
INTR 48144 ... 48440 OCKHAMG-CDS
INTR 48542 ... 48757 OCKHAMG-CDS

INTR 51354 ... 51417 OCKHAMG-CDNA

INTR 48908 ... 49027 OCKHAMG-CDS INTR 49117 ... 49654 OCKHAMG-CDS

TERM 50075 ... 50184 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 372
- Ceres seq id 2025475
- Alternative transcription start site(s) located in SEQ ID NO 372:

1242,1251,1270,1271,1275,1276,1280,1287,1288,1290,1292,1293,1 301,1302,1305,1307,1314,1318,1321,1325

35 1326,1330

- Clone 27681 starts at 1242 and ends at 2029 in cDNA.
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 373
  - Ceres seq id 2025476
  - Location of start within SEQ ID NO 372: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 245
    - qi No. 2739366
    - Description: (AC002505) SF16 like protein

[Arabidopsis thaliana]

- 50 % Identity: 45
  - Alignment Length: 373

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234
             - Location of Alignment in SEQ ID NO 373: from 61
to 418
    (B) Polypeptide Sequence
         - Pat. Appln. SEQ ID NO 374
         - Ceres seq id 2025477
         - Location of start within SEQ ID NO 372: at 316 nt.
        (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
        (Dp) Related Amino Acid Sequences
             - Alignment No. 246
             - qi No. 2739366
             - Description: (AC002505) SF16 like protein
[Arabidopsis thaliana]
             - % Identity: 45
             - Alignment Length: 373
             - Location of Alignment in SEQ ID NO 374: from 1 to
313
    (B) Polypeptide Sequence
         - Pat. Appln. SEQ ID NO 375
         - Ceres seq id 2025478
         - Location of start within SEQ ID NO 372: at 364 nt.
         - Location of Signal Peptide Cleavage Site within SEQ
ID NO 375: at 21 aa.
        (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
        (Dp) Related Amino Acid Sequences
             - Alignment No. 247
             - qi No. 2739366
             - Description: (AC002505) SF16 like protein
[Arabidopsis thaliana]
            - % Identity:
                             45
            - Alignment Length: 373
            - Location of Alignment in SEQ ID NO 375: from 1 to
297
Maximum Length Sequence:
 related to:
Clone IDs:
       14564
Public Genomic DNA:
       gi No: 4415905
       Predicted Exons:

      INTR
      3235
      ...
      3181
      OCKHAMG-CDNA

      INTR
      2694
      ...
      2619
      OCKHAMG-CDNA

      INTR
      2509
      ...
      2408
      OCKHAMG-CDNA

      INTR
      2319
      ...
      2171
      OCKHAMG-CDNA
```

2076 ... 1784 OCKHAMG-CDNA

235

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3191 ... 3181
                                        OCKHAMG-CDS
              INIT
                                2619 OCKHAMG-CDS
                     2694 ...
              INTR
                     2509 ... 2408
              INTR
                                       OCKHAMG-CDS
 5
              INTR
                     2319
                          ... 2171 OCKHAMG-CDS
                          ... 1920
              TERM 2076
                                       OCKHAMG-CDS
           qi No: 4581138
           Predicted Exons:
                     100911... 100857 OCKHAMG-CDNA
              INTR
10
              INTR 100370... 100295 OCKHAMG-CDNA
              INTR 100185... 100084 OCKHAMG-CDNA INTR 99995 ... 99847 OCKHAMG-CDNA
              INTR 99752 ... 99460 OCKHAMG-CDNA
15
              INIT 100867... 100857 OCKHAMG-CDS
              INTR 100370... 100295 OCKHAMG-CDS
              INTR 100185... 100084 OCKHAMG-CDS
              INTR 99995 ... 99847
                                        OCKHAMG-CDS
                   99752 ... 99596 OCKHAMG-CDS
              TERM
     (Ac) cDNA Polynucleotide Sequence
20
          - Pat. Appln. SEQ ID NO 376
          - Ceres seq id 2025524
          - Alternative transcription start site(s) located in SEQ
     ID NO 376:
25
            -35, -26, -25, -2, 2, 3, 4, 5, 6, 7, 8, 9, 15, 16, 17, 32
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 377
            - Ceres seq id 2025525
            - Location of start within SEQ ID NO 376: at 47 nt.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Ribosomal protein L24e
35
               - Location within SEQ ID NO 377: from 3 to 73 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 248
               - qi No. 6094040
40
               - Description: 60S RIBOSOMAL PROTEIN L24
     >qi|2961300|emb|CAA12358| (AJ225027) ribosomal protein L24
     [Cicer arietinum]
               - % Identity: 88.3
               - Alignment Length: 163
45
               - Location of Alignment in SEQ ID NO 377: from 1 to
     163
        (B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 378
- Ceres seq id 2025526

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- Location of start within SEQ ID NO 376: at 212 nt.

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
 5
               - Alignment No. 249
               - qi No. 6094040
               - Description: 60S RIBOSOMAL PROTEIN L24
     >qi|2961300|emb|CAA12358| (AJ225027) ribosomal protein L24
     [Cicer arietinum]
10
               - % Identity: 88.3
               - Alignment Length: 163
               - Location of Alignment in SEQ ID NO 378: from 1 to
     108
15
     Maximum Length Sequence:
      related to:
     Clone IDs:
           20767
     Public Genomic DNA:
20
           qi No: 4519195
           Predicted Exons:
              INIT
                     50757 ... 50625 OCKHAMG-CDS
              INTR 50001 ... 49847 OCKHAMG-CDS
              INTR 49644 ... 49561 OCKHAMG-CDS INTR 49391 ... 49236 OCKHAMG-CDS
25
                     49128 ... 49023 OCKHAMG-CDS
              INTR
              TERM 48810 ... 48662 CCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 379
30
          - Ceres seq id 2025544
          - Alternative transcription start site(s) located in SEQ
     ID NO 379:
     656, 672, 680, 685, 701, 703, 706, 709, 713, 717, 728, 736, 741, 743
35
          - Clone 20767 starts at 656 and ends at 1623 in cDNA.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 380
            - Ceres seq id 2025545
40
            - Location of start within SEQ ID NO 379: at 599 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Domain of unknown function
45
               - Location within SEQ ID NO 380: from 134 to 247
     aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 250
50
               - qi No. 1708292
```

237 - Description: HEAT-RESPONSIVE PROTEIN 12 >gi|1255116 (U50631) heat-responsive protein [Mus musculus] - % Identity: 54.3 - Alignment Length: 129 5 - Location of Alignment in SEQ ID NO 380: from 119 to 247 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 381 10 - Ceres seq id 2025546 - Location of start within SEQ ID NO 379: at 1 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 381: at 50 aa. 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Zinc finger, C3HC4 type (RING finger) - Location within SEQ ID NO 381: from 171 to 211 aa. 20 (Dp) Related Amino Acid Sequences - Alignment No. 251 - qi No. 2435518 - Description: (AF024504) contains similarity to C3HC4-type zinc fingers [Arabidopsis thaliana] 25 - % Identity: 31.4 - Alignment Length: 221 - Location of Alignment in SEQ ID NO 381: from 35 to 211 30 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 382 - Ceres seq id 2025547 - Location of start within SEQ ID NO 379: at 16 nt. - Location of Signal Peptide Cleavage Site within SEQ 35 ID NO 382: at 45 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Zinc finger, C3HC4 type (RING finger) 40 - Location within SEQ ID NO 382: from 166 to 206 aa. (Dp) Related Amino Acid Sequences - Alignment No. 252 45 - gi No. 2435518 - Description: (AF024504) contains similarity to C3HC4-type zinc fingers [Arabidopsis thaliana] - % Identity: 31.4

- Alignment Length: 221

238

- Location of Alignment in SEQ ID NO 382: from 30 to 206

Maximum Length Sequence:

related to:

Clone IDs:

7802

Public Genomic DNA:

qi No: 4415920

10 Predicted Exons:

INTR 60672 ... 61210 OCKHAMG-CDNA

SINGLE 60712 ... 61119 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 383

- Ceres seq id 2026207

- Alternative transcription start site(s) located in SEQ ID NO 383:

2,3,4,5

20

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 384
  - Ceres seq id 2026208
  - Location of start within SEQ ID NO 383: at 2 nt.

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- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 253
  - gi No. 4757974
- Description: ref|NP\_004057.1|pCETN1| centrin, EFhand protein, 1 >gi|2493440|sp|Q12798|CAT2\_HUMAN CALTRACTIN, ISOFORM 2 (CENTRIN) >gi|414993 (U03270) centrin [Homo sapiens]
  - % Identity: 34.3
  - Alignment Length: 102
  - Location of Alignment in SEQ ID NO 384: from 19

to 120

- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 385
  - Ceres seq id 2026209
  - Location of start within SEQ ID NO 383: at 41 nt.
- 45 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 254
    - gi No. 4757974
- Description: ref[NP\_004057.1|pCETN1| centrin, EF-hand protein, 1 >gi|2493440|sp|Q12798|CAT2 HUMAN CALTRACTIN,

239

ISOFORM 2 (CENTRIN) >gi|414993 (U03270) centrin [Homo
sapiens]

- % Identity: 34.3
- Alignment Length: 102
- 5 Location of Alignment in SEQ ID NO 385: from 6 to 107
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 386
- 10 Ceres seg id 2026210
  - Location of start within SEQ ID NO 383: at 62 nt.
  - - (Dp) Related Amino Acid Sequences
      - Alignment No. 255
      - qi No. 4757974
  - Description: ref|NP\_004057.1|pCETN1| centrin, EF-hand protein, 1 >qi|2493440|sp|Q12798|CAT2 HUMAN CALTRACTIN,
- 20 ISOFORM 2 (CENTRIN) >gi|414993 (U03270) centrin [Homo sapiens]
  - % Identity: 34.3
  - Alignment Length: 102
  - Location of Alignment in SEQ ID NO 386: from 1 to

25 100

15

Maximum Length Sequence:

Public Genomic DNA:

gi No: 4038029

30 Predicted Exons:

INIT	46188	 46125	GENBANK
TERM	46021	 45852	GENBANK

INTR 46258 ... 46125 OCKHAMG-CDNA 35 INTR 46021 ... 45663 OCKHAMG-CDNA

> INIT 47157 ... 47119 OCKHAMG-CDS INTR 46278 ... 46125 OCKHAMG-CDS TERM 46021 ... 45852 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 387
- Ceres seq id 2026982

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 388
  - Ceres seq id 2026983
  - Location of start within SEQ ID NO 387: at 1 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 388: at 73 aa.

240

5	<pre>(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)</pre>
10	<pre>(Dp) Related Amino Acid Sequences</pre>
20	(B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 389 - Ceres seq_id 2026984 - Location of start within SEQ ID NO 387: at 34 nt Location of Signal Peptide Cleavage Site within SEQ ID NO 389: at 54 aa.
25	<pre>(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)</pre>
30	(Dp) Related Amino Acid Sequences - Alignment No. 257 - gi No. 1209258 - Description: (L31937) protease inhibitor II
35	[Brassica rapa] - % Identity: 87 - Alignment Length: 77 - Location of Alignment in SEQ ID NO 389: from 33 to 109
40	<ul><li>(B) Polypeptide Sequence</li><li>- Pat. Appln. SEQ ID NO 390</li><li>- Ceres seq_id 2026985</li><li>- Location of start within SEQ ID NO 387: at 130 nt.</li></ul>
	- Location of Signal Peptide Cleavage Site within SEQ ID NO 390: at 22 aa.
45	<pre>(C) Nomination and Annotation of Domains within Predicted Polypeptide(s)</pre>
50	

(Dp) Related Amino Acid Sequences

WO 00/40695 PCT/US00/00466 241 - Alignment No. 258 - qi No. 1209258 - Description: (L31937) protease inhibitor II

[Brassica rapa]

- % Identity: 87

- Alignment Length: 77

- Location of Alignment in SEQ ID NO 390: from 1 to

77

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 391
  - Ceres seq id 2026986
  - Location of start within SEQ ID NO 387: at 30 nt.
- 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences

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Maximum Length Sequence:
```

20 related to:

Clone IDs:

251127

Public Genomic DNA:

qi No: 4220640

25 Predicted Exons:

> 67833 ... 68135 OCKHAMG-CDS INIT 68253 ... 68366 OCKHAMG-CDS INTR INTR 68457 ... 68552 OCKHAMG-CDS INTR 68720 ... 68794 OCKHAMG-CDS INTR 68898 ... 68965 OCKHAMG-CDS INTR 69056 ... 69066 OCKHAMG-CDS INTR 69422 ... 69624 OCKHAMG-CDS INTR 69858 ... 70202 OCKHAMG-CDS INTR 70315 ... 70428 OCKHAMG-CDS INTR 70516 ... 70611 OCKHAMG-CDS INTR 70923 ... 70997 OCKHAMG-CDS INTR 71103 ... 71173 OCKHAMG-CDS INTR 71737 ... 71749 OCKHAMG-CDS TERM

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 392
- Ceres seq id 2027300
- Alternative transcription start site(s) located in SEQ ID NO 392:

38,53,55,63,82,84,85,103,160

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 393
  - Ceres seq id 2027301
  - Location of start within SEQ ID NO 392: at 177 nt.

242

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(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 259
5
               - qi No. 5020103
               - Description: (AF148219) fibrillin [Nostoc
     PCC80091
               - % Identity: 30.3
               - Alignment Length: 122
10
               - Location of Alignment in SEQ ID NO 393: from 75
    to 189
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 394
15
            - Ceres seq id 2027302
            - Location of start within SEQ ID NO 392: at 213 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
20
           (Dp) Related Amino Acid Sequences
               - Alignment No. 260
               - gi No. 5020103
               - Description: (AF148219) fibrillin [Nostoc
     PCC80091
25
               - % Identity: 30.3
               - Alignment Length: 122
               - Location of Alignment in SEQ ID NO 394: from 63
     to 177
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 395
            - Ceres seq id 2027303
            - Location of start within SEQ ID NO 392: at 252 nt.
35
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 261
               - qi No. 5020103
40
               - Description: (AF148219) fibrillin [Nostoc
     PCC80091
               - % Identity: 30.3
               - Alignment Length: 122
               - Location of Alignment in SEQ ID NO 395: from 50
45
    to 164
    Maximum Length Sequence:
      related to:
     Clone IDs:
50
           271260
```

Public Genomic DNA:

243 qi No: 6056186 Predicted Exons: INIT 66600 ... 66465 OCKHAMG-CDS TERM 66390 ... 66083 OCKHAMG-CDS 5 gi No: 6087919 Predicted Exons: 31657 ... 31522 OCKHAMG-CDS INIT TERM 31447 ... 31140 OCKHAMG-CDS qi No: 6119514 10 Predicted Exons: 32265 ... 32130 OCKHAMG-CDS INIT 32055 ... 31748 OCKHAMG-CDS TERM qi No: 6143857 Predicted Exons: 34633 ... 34768 OCKHAMG-CDS 15 INIT 34843 ... 35150 TERM OCKHAMG-CDS qi No: 6175136 Predicted Exons: INIT 34136 ... 34271 OCKHAMG-CDS 20 TERM 34346 ... 34653 OCKHAMG-CDS qi No: 6223634 Predicted Exons: 34137 ... 34272 INIT OCKHAMG-CDS 34347 ... 34654 TERM OCKHAMG-CDS 25 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 396 - Ceres seq id 2027375 30 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 397 - Ceres seq id 2027376 - Location of start within SEQ ID NO 396: at 2 nt. 35 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Bacterial mutT protein - Location within SEQ ID NO 397: from 27 to 68 aa. 40 (Dp) Related Amino Acid Sequences - Alignment No. 262 - qi No. 2622420 - Description: (AE000895) mutator MutT protein [Methanobacterium thermoautotrophicum] 45 - % Identity: 42.3 - Alignment Length: 98 - Location of Alignment in SEQ ID NO 397: from 13 to 109 50 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 398

244

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- Ceres seq id 2027377
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- Location of start within SEQ ID NO 396: at 17 nt.
- (C) Nomination and Annotation of Domains within
  Predicted Polypeptide(s)
  - Bacterial mutT protein
  - Location within SEQ ID NO 398: from 22 to 63 aa.
  - (Dp) Related Amino Acid Sequences
- Alignment No. 263
  - gi No. 2622420
  - Description: (AE000895) mutator MutT protein [Methanobacterium thermoautotrophicum]
    - % Identity: 42.3
- Alignment Length: 98
  - Location of Alignment in SEQ ID NO 398: from 8 to 104
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 399
      - Ceres seq id 2027378
      - Location of start within SEQ ID NO 396: at 188 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 264
    - gi No. 2622420
    - Description: (AE000895) mutator MutT protein
- 30 [Methanobacterium thermoautotrophicum]
  - % Identity: 42.3
  - Alignment Length: 98
  - Location of Alignment in SEQ ID NO 399: from 1 to

47

35

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Maximum Length Sequence:

Public Genomic DNA:

gi No: 5868932

Predicted Exons:

40 INIT 99857 ... 96712 GENBANK INTR 96123 ... 95883 GENBANK TERM 95642 ... 95199 GENBANK

> INTR 96097 ... 95883 OCKHAMG-CDNA INTR 95642 ... 95047 OCKHAMG-CDNA

> > INIT 96035 ... 95883 OCKHAMG-CDS TERM 95642 ... 95199 OCKHAMG-CDS

50 (Ac) cDNA Polynucleotide Sequence
- Pat. Appln. SEQ ID NO 400

245

- Ceres seq id 2028729
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 401
  - Ceres seq id 2028730
  - Location of start within SEQ ID NO 400: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - C2 domain
  - Location within SEQ ID NO 401: from 671 to 762 aa.
- 15 (Dp) Related Amino Acid Sequences
  - Alignment No. 265
  - qi No. 3142295
  - Description: (AC002411) Strong similarity to phosphoribosylanthranilate transferase gb/D86180 from Pisum sativum. [Arabidopsis thaliana]
    - % Identity: 45.6
    - Alignment Length: 865
    - Location of Alignment in SEQ ID NO 401: from 211 to 1050  $\,$

25

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 402
  - Ceres seq id 2028731
  - Location of start within SEQ ID NO 400: at 199 nt.

30

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
  - C2 domain
  - Location within SEQ ID NO 402: from 605 to 696
- 35 aa.
- (Dp) Related Amino Acid Sequences
  - Alignment No. 266
  - gi No. 3142295
- Description: (AC002411) Strong similarity to phosphoribosylanthranilate transferase gb/D86180 from Pisum sativum. [Arabidopsis thaliana]
  - % Identity: 45.6
  - Alignment Length: 865
- Location of Alignment in SEQ ID NO 402: from 145 to 984
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 403
    - Ceres seq id 2028732
    - Location of start within SEQ ID NO 400: at 232 nt.

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246

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(C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
          - C2 domain
          - Location within SEQ ID NO 403: from 594 to 685
aa.
      (Dp) Related Amino Acid Sequences
        - Alignment No. 267
          - qi No. 3142295
          - Description: (AC002411) Strong similarity to
phosphoribosylanthranilate transferase gb|D86180 from Pisum
sativum. [Arabidopsis thaliana]
          - % Identity:
          - Alignment Length: 865
          - Location of Alignment in SEQ ID NO 403: from 134
to 973
Maximum Length Sequence:
 related to:
Clone IDs:
      9214
(Ac) cDNA Polynucleotide Sequence
     - Pat. Appln. SEQ ID NO 404
     - Ceres seq id 2029079
     - Alternative transcription start site(s) located in SEO
ID NO 404:
       -4, -1, 2, 3, 4, 8, 9, 10, 11, 14, 15, 16, 17, 18, 20, 23, 24
   (B) Polypeptide Sequence
       - Pat. Appln. SEQ ID NO 405
       - Ceres seq id 2029080
       - Location of start within SEQ ID NO 404: at 3 nt.
      (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
      (Dp) Related Amino Acid Sequences
          - Alignment No. 268
          - qi No. 132825
          - Description: 50S RIBOSOMAL PROTEIN CL25,
CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein
PsCL25 precursor, chloroplast - garden pea
>gi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein
(AA -30 to 74) [Pisum sativum]
          - % Identity:
                         56.1
          - Alignment Length: 82
          - Location of Alignment in SEQ ID NO 405: from 14
to 95
```

50 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 406

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247
            - Ceres seq id 2029081
            - Location of start within SEQ ID NO 404: at 39 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 406: at 16 aa.
5
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 269
10
               - qi No. 132825
               - Description: 50S RIBOSOMAL PROTEIN CL25,
     CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein
     PsCL25 precursor, chloroplast - garden pea
     >qi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein
15
     (AA - 30 \text{ to } 74) [Pisum sativum]
               - % Identity: 56.1
               - Alignment Length: 82
               - Location of Alignment in SEQ ID NO 406: from 2 to
     83
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 407
            - Ceres seq id 2029082
            - Location of start within SEQ ID NO 404: at 144 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
                - Alignment No. 270
30
               - qi No. 132825
               - Description: 50S RIBOSOMAL PROTEIN CL25,
     CHLOROPLAST PRECURSOR >gi|71308|pir||R5PM25 ribosomal protein
     PsCL25 precursor, chloroplast - garden pea
     >qi|20877|emb|CAA32187| (X14022) PsCL25 ribosomal preprotein
35
     (AA - 30 \text{ to } 74) [Pisum sativum]
               - % Identity: 56.1
               - Alignment Length: 82
               - Location of Alignment in SEQ ID NO 407: from 1 to
     48
40
     Maximum Length Sequence:
     Public Genomic DNA:
           gi No: 6434227
           Predicted Exons:
45
               INTR 50076 ... 50370 OCKHAMG-CDNA
              INTR 50793 ... 50874 OCKHAMG-CDNA INTR 51163 ... 51227 OCKHAMG-CDNA
```

INTR 51331 ... 51417

50

INTR 51529 ... 51786 OCKHAMG-CDNA

INIT 50200 ... 50370 OCKHAMG-CDS

OCKHAMG-CDNA

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248
                                50874 OCKHAMG-CDS
              INTR
                     50793 ...
                     51163 ... 51227 OCKHAMG-CDS
              INTR
                     51331 ... 51417 OCKHAMG-CDS
              INTR
              TERM 51529 ... 51630 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 408
          - Ceres seq id 2029225
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 409
            - Ceres seq id 2029226
            - Location of start within SEQ ID NO 408: at 125 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 271
20
               - qi No. 1352051
               - Description: ATP SYNTHASE D CHAIN, MITOCHONDRIAL
     >qi|484646|pir||JS0739 H+-transporting ATP synthase (EC
     3.6.1.34) chain d precursor - rat >gi|286204|dbj|BAA02422|
     (D13120) ATP synthase subunit d precursor [Rattus norvegicus]
               - % Identity: 28.6
25
               - Alignment Length: 145
               - Location of Alignment in SEQ ID NO 409: from 15
     to 154
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 410
            - Ceres seq id 2029227
            - Location of start within SEQ ID NO 408: at 194 nt.
35
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 272
               - qi No. 1352051
40
               - Description: ATP SYNTHASE D CHAIN, MITOCHONDRIAL
     >qi|484646|pir||JS0739 H+-transporting ATP synthase (EC
     3.6.1.34) chain d precursor - rat >gi|286204|dbj|BAA02422|
     (D13120) ATP synthase subunit d precursor [Rattus norvegicus]
               - % Identity:
                              28.6
               - Alignment Length: 145
45
               - Location of Alignment in SEQ ID NO 410: from 1 to
     131
     Maximum Length Sequence:
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Public Genomic DNA:

gi No: 6056185

249 Predicted Exons: 21555 ... 22692 OCKHAMG-CDNA INTR 22891 ... 23339 INTR OCKHAMG-CDNA 5 INTR 22502 ... 22692 OCKHAMG-CDNA 23338 OCKHAMG-CDNA INTR 22891 ... 21706 ... 22692 OCKHAMG-CDS INIT 23124 OCKHAMG-CDS TERM 22891 ... gi No: 6091703 10 Predicted Exons: INTR 21555 ... 22692 OCKHAMG-CDNA 22891 ... 23339 INTR OCKHAMG-CDNA 15 INTR 22502 ... 22692 OCKHAMG-CDNA 22891 ... 23338 OCKHAMG-CDNA INTR 21706 ... 22692 OCKHAMG-CDS INIT 22891 ... 23124 OCKHAMG-CDS TERM 20 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 411 - Ceres seq id 2029422 25 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 412 - Ceres seq id 2029423 - Location of start within SEQ ID NO 411: at 152 nt. 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - CONSTANS family zinc finger - Location within SEQ ID NO 412: from 17 to 57 aa. 35 (Dp) Related Amino Acid Sequences - Alignment No. 273 - qi No. 1076301 - Description: CONSTANS protein - Arabidopsis thaliana >qi|1161514|emb|CAA64407| (X94937) CONSTANS protein 40 [Arabidopsis thaliana] - % Identity: 49.2 - Alignment Length: 63 - Location of Alignment in SEQ ID NO 412: from 12 45 to 74 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 413 - Ceres seq id 2029424 - Location of start within SEQ ID NO 411: at 155 nt. 50

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250
      (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
          - CONSTANS family zinc finger
          - Location within SEQ ID NO 413: from 16 to 56 aa.
      (Dp) Related Amino Acid Sequences
          - Alignment No. 274
          - qi No. 1076301
          - Description: CONSTANS protein - Arabidopsis
thaliana >qi|1161514|emb|CAA64407| (X94937) CONSTANS protein
[Arabidopsis thaliana]
          - % Identity:
                        49.2
          - Alignment Length: 63
          - Location of Alignment in SEQ ID NO 413: from 11
to 73
   (B) Polypeptide Sequence
       - Pat. Appln. SEQ ID NO 414
       - Ceres seq id 2029425
       - Location of start within SEQ ID NO 411: at 443 nt.
      (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
      (Dp) Related Amino Acid Sequences
          - Alignment No. 275
          - qi No. 1076301
          - Description: CONSTANS protein - Arabidopsis
thaliana >qi|1161514|emb|CAA64407| (X94937) CONSTANS protein
[Arabidopsis thaliana]
          - % Identity: 69.8
          - Alignment Length: 43
          - Location of Alignment in SEQ ID NO 414: from 260
to 302
Maximum Length Sequence:
Public Genomic DNA:
      gi No: 5832738
      Predicted Exons:
                           56150
         INTR
               56241 ...
                                   OCKHAMG-CDNA
                56041 ... 55864
                                   OCKHAMG~CDNA
         INTR
                          55317
         INTR
               55493 ...
                                   OCKHAMG-CDNA
               55216 ...
                          55085
                                   OCKHAMG~CDNA
         INTR
                          54803
               54898 ...
         INTR
                                   OCKHAMG-CDNA
               54725 ... 54609 OCKHAMG-CDNA
         INTR
               54510 ... 54415 OCKHAMG-CDNA
         INTR
               54288 ... 54176 OCKHAMG-CDNA
         INTR
                          53779
         INTR
               54077 ...
                                   OCKHAMG-CDNA
```

56181 ... 56150 OCKHAMG-CDS

OCKHAMG-CDS

OCKHAMG-CDS

56041 ... 55864

55493 ... 55317

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INIT

INTR

INTR

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251
                                55085 OCKHAMG-CDS
                    55216 ...
              INTR
                               54803 OCKHAMG-CDS
              INTR
                    54898 ...
                    54725 ... 54609 OCKHAMG-CDS
              INTR
                    54510 ... 54415 OCKHAMG-CDS
              INTR
                    54288 ... 54176 OCKHAMG-CDS
5
              INTR
                    54077 ... 53954 OCKHAMG-CDS
             TERM
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 415
10
          - Ceres seq id 2029806
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 416
            - Ceres seq id 2029807
15
            - Location of start within SEQ ID NO 415: at 1 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Aminotransferase class IV
20
               - Location within SEQ ID NO 416: from 68 to 352 aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 276
               - qi No. 3540183
25
               - Description: (AC004122) Highly Similar to
     branched-chain amino acid aminotransferase [Arabidopsis
     thalianal
               - % Identity: 53.8
               - Alignment Length: 318
30
               - Location of Alignment in SEQ ID NO 416: from 55
     to 371
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 417
35
            - Ceres seq id 2029808
            - Location of start within SEQ ID NO 415: at 61 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
40
               - Aminotransferase class IV
               - Location within SEO ID NO 417: from 48 to 332 aa.
           (Dp) Related Amino Acid Sequences
45
               - Alignment No. 277
               - qi No. 3540183
               - Description: (AC004122) Highly Similar to
     branched-chain amino acid aminotransferase [Arabidopsis
     thaliana
50
               - % Identity: 53.8
```

- Alignment Length: 318

252

- Location of Alignment in SEQ ID NO 417: from 35 to 351  $\,$ 

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 418
  - Ceres seq id 2029809
  - Location of start within SEQ ID NO 415: at 163 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Aminotransferase class IV
  - Location within SEQ ID NO 418: from 14 to 298 aa.
  - (Dp) Related Amino Acid Sequences
- Alignment No. 278
  - qi No. 3540183
  - Description: (AC004122) Highly Similar to branched-chain amino acid aminotransferase [Arabidopsis thaliana]
- 20 % Identity: 53.8
  - Alignment Length: 318
  - Location of Alignment in SEQ ID NO 418: from 1 to

317

25 Maximum Length Sequence:

Public Genomic DNA:

gi No: 5822667

Predicted Exons:

INTR 59013 ... 59999 OCKHAMG-CDNA

30

5

SINGLE 59071 ... 59769 OCKHAMG-CDS

gi No: 6041831

Predicted Exons:

INTR 53943 ... 54929 OCKHAMG-CDNA

35

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SINGLE 54001 ... 54699 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 419
- 40 Ceres seq id 2030038
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 420
    - Ceres seq id 2030039
    - Location of start within SEQ ID NO 419: at 2 nt.
  - (C) Nomination and Annotation of Domains within  $Predicted\ Polypeptide(s)$ 
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 279

253

- qi No. 4009495

- Description: (AF068904) YlmG [Staphylococcus

aureus]

- % Identity: 31.7

- Alignment Length: 63

- Location of Alignment in SEQ ID NO 420; from 177

to 239

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 421
  - Ceres seq id 2030040
  - Location of start within SEQ ID NO 419: at 59 nt.
- Location of Signal Peptide Cleavage Site within SEQ

ID NO 421: at 19 aa.

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 280
    - qi No. 4009495
    - Description: (AF068904) YlmG [Staphylococcus

aureus]

- % Identity: 31.7
- Alignment Length: 63
- 25 Location of Alignment in SEQ ID NO 421: from 158

to 220

Maximum Length Sequence:

Public Genomic DNA:

30 gi No: 4539290 Predicted Exons:

INIT	6134	 6132	GENBANK
INTR	6021	 5973	GENBANK
INTR	5588	 5309	GENBANK
INTR	5226	 5106	GENBANK
TERM	4951	 4871	GENBANK

INTR 6208 ... 6132 OCKHAMG-CDNA
INTR 6021 ... 5973 OCKHAMG-CDNA
40 INTR 5588 ... 5309 OCKHAMG-CDNA
INTR 5226 ... 5106 OCKHAMG-CDNA
INTR 4951 ... 4724 OCKHAMG-CDNA

INIT 6134 ... 6132 OCKHAMG-CDS

INTR 6021 ... 5973 OCKHAMG-CDS

INTR 5588 ... 5309 OCKHAMG-CDS

INTR 5226 ... 5106 OCKHAMG-CDS

TERM 4951 ... 4871 OCKHAMG-CDS

gi No: 4914454 Predicted Exons:

INTR 39557 ... 39481 OCKHAMG-CDNA

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162

Predicted Polypeptide(s)

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254
                39370 ...
                           39322 OCKHAMG-CDNA
         INTR
                38937 ... 38658 OCKHAMG-CDNA
38575 ... 38455 OCKHAMG-CDNA
         INTR
         INTR
                38300 ... 38073 OCKHAMG-CDNA
         INTR
(Ac) cDNA Polynucleotide Sequence
     - Pat. Appln. SEQ ID NO 422
     - Ceres seq id 2031778
   (B) Polypeptide Sequence
       - Pat. Appln. SEQ ID NO 423
       - Ceres seq id 2031779
       - Location of start within SEQ ID NO 422: at 3 nt.
      (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
      (Dp) Related Amino Acid Sequences
          - Alignment No. 281
          - qi No. 3688432
          - Description: (AJ011705) 40S ribosomal protein S10
[Lumbricus rubellus]
          - % Identity: 53.1
          - Alignment Length: 164
          - Location of Alignment in SEQ ID NO 423: from 25
to 186
   (B) Polypeptide Sequence
       - Pat. Appln. SEQ ID NO 424
       - Ceres seq id 2031780
       - Location of start within SEQ ID NO 422: at 75 nt.
      (C) Nomination and Annotation of Domains within
Predicted Polypeptide(s)
      (Dp) Related Amino Acid Sequences
          - Alignment No. 282
          - qi No. 3688432
          - Description: (AJ011705) 40S ribosomal protein S10
[Lumbricus rubellus]
          - % Identity:
                          53.1
          - Alignment Length: 164
          - Location of Alignment in SEQ ID NO 424: from 1 to
   (B) Polypeptide Sequence
       - Pat. Appln. SEQ ID NO 425
       - Ceres seq id 2031781
       - Location of start within SEQ ID NO 422: at 210 nt.
```

(C) Nomination and Annotation of Domains within

255 (Dp) Related Amino Acid Sequences - Alignment No. 283 - qi No. 3688432 - Description: (AJ011705) 40S ribosomal protein S10 5 [Lumbricus rubellus] - % Identity: 53.1 - Alignment Length: 164 - Location of Alignment in SEQ ID NO 425: from 1 to 117 10 Maximum Length Sequence: Public Genomic DNA: gi No: 6434226 Predicted Exons: 15 INTR 17873 ... 17514 OCKHAMG-CDNA INTR 17046 ... 16869 OCKHAMG-CDNA INIT 17854 ... 17514 OCKHAMG-CDS TERM 17046 ... 17028 OCKHAMG-CDS 20 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 426 - Ceres seq id 2032723 25 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 427 - Ceres seq id 2032724 - Location of start within SEQ ID NO 426: at 20 nt. - Location of Signal Peptide Cleavage Site within SEQ 30 ID NO 427: at 24 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Plant lipid transfer protein family 35 - Location within SEQ ID NO 427: from 28 to 115 aa. (Dp) Related Amino Acid Sequences - Alignment No. 284 - gi No. 899224 40 - Description: (X60318) E2 [Brassica napus] >qi|384340|prf||1905428A phospholipid transfer protein [Brassica napus] - % Identity: 79 45 - Alignment Length: 119 - Location of Alignment in SEQ ID NO 427: from 1 to 119 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 428

- Ceres seq id 2032725

256

- Location of start within SEQ ID NO 426: at 71 nt.
- - Plant lipid transfer protein family
  - Location within SEQ ID NO 428: from 11 to 98 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 285
    - gi No. 899224
- Description: (X60318) E2 [Brassica napus] >qi|384340|prf||1905428A phospholipid transfer protein

[Brassica napus]

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- % Identity: 79
- Alignment Length: 119
- Location of Alignment in SEQ ID NO 428: from 1 to 102
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 429
  - Ceres seq id 2032726
  - Location of start within SEQ ID NO 426: at 1 nt.
- (C) Nomination and Annotation of Domains within 25 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences

Maximum Length Sequence:

Public Genomic DNA:

30 qi No: 6143856

Predicted Exons:

INIT 30571 ... 30464 OCKHAMG-CDS TERM 30327 ... 30238 OCKHAMG-CDS

qi No: 6223633

35 Predicted Exons:

INIT 30571 ... 30464 OCKHAMG-CDS TERM 30327 ... 30238 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 430
    - Ceres seq id 2035536
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 431
    - Ceres seq id 2035537
    - Location of start within SEQ ID NO 430: at 1 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 431: at 20 aa.

50

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257

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(C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 286
 5
               - qi No. 3024192
               - Description: NADH-UBIQUINONE OXIDOREDUCTASE MWFE
    SUBUNIT (COMPLEX I-MWFE) (CI-MWFE) >gi|2275037|emb|CAA68977|
     (Y07708) NADH dehydrogenase [Mus musculus]
             - % Identity: 31.5
10
               - Alignment Length: 54
               - Location of Alignment in SEQ ID NO 431: from 4 to
    56
        (B) Polypeptide Sequence
15
            - Pat. Appln. SEQ ID NO 432
            - Ceres seq id 2035538
            - Location of start within SEQ ID NO 430: at 2 nt.
           (C) Nomination and Annotation of Domains within
20
    Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 433
25
            - Ceres seq id 2035539
            - Location of start within SEQ ID NO 430: at 25 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 433: at 17 aa.
30
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 287
               - qi No. 3024192
35
               - Description: NADH-UBIQUINONE OXIDOREDUCTASE MWFE
     SUBUNIT (COMPLEX I-MWFE) (CI-MWFE) >qi|2275037|emb|CAA68977|
     (Y07708) NADH dehydrogenase [Mus musculus]
               - % Identity: 31.5
               - Alignment Length: 54
               - Location of Alignment in SEQ ID NO 433: from 1 to
40
     48
    Maximum Length Sequence:
     Public Genomic DNA:
45
           gi No: 6136357
           Predicted Exons:
              INIT 51922 ... 51830 OCKHAMG-CDS
              INTR 51129 ... 51055 OCKHAMG-CDS
              INTR 50958 ... 50806 OCKHAMG-CDS
50
              INTR 50706 ... 50608 OCKHAMG-CDS
```

TERM 50326 ... 50243 OCKHAMG-CDS

```
258
           qi No: 6143858
           Predicted Exons:
                     51922 ...
                                 51830 OCKHAMG-CDS
               INIT
                     51129 ...
                                 51055 OCKHAMG-CDS
              INTR
              INTR 50958 ... 50806 OCKHAMG-CDS INTR 50706 ... 50608 OCKHAMG-CDS
 5
              TERM 50326 ... 50243 OCKHAMG-CDS
           qi No: 6449510
           Predicted Exons:
                                 9111 OCKHAMG-CDNA
9360 OCKHAMG-CDNA
10
              INTR
                     9035 ...
                     9208 ... 9360
              INTR
              INTR 9460 ... 9558 OCKHAMG-CDNA INTR 9840 ... 9992 OCKHAMG-CDNA
15
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 434
          - Ceres seq id 2035575
20
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 435
            - Ceres seq id 2035576
            - Location of start within SEQ ID NO 434: at 1 nt.
25
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
                - Alignment No. 288
               - qi No. 6066609
               - Description: (AJ009987) chloroplast channel
30
     forming outer membrane protein [Pisum sativum]
               - % Identity: 54.2
                - Alignment Length: 177
               - Location of Alignment in SEQ ID NO 435: from 1 to
35
     167
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 436
            - Ceres seq id 2035577
40
            - Location of start within SEQ ID NO 434: at 13 nt.
            (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
            (Dp) Related Amino Acid Sequences
                - Alignment No. 289
45
               - qi No. 6066609
                - Description: (AJ009987) chloroplast channel
     forming outer membrane protein [Pisum sativum]
               - % Identity: 54.2
```

- Alignment Length: 177

259

- Location of Alignment in SEQ ID NO 436: from 1 to 163

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 437
  - Ceres seq id 2035578
  - Location of start within SEQ ID NO 434: at 40 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 290
    - gi No. 6066609
    - Description: (AJ009987) chloroplast channel
- forming outer membrane protein [Pisum sativum]
  - % Identity: 54.2
  - Alignment Length: 177
  - Location of Alignment in SEQ ID NO 437: from 1 to

154

Maximum Length Sequence:

Public Genomic DNA:

gi No: 6434213

Predicted Exons:

25 INIT 54056 ... 54085 OCKHAMG-CDS INTR 54178 ... 54325 OCKHAMG-CDS INTR 54401 ... 54572 OCKHAMG-CDS INTR 54807 ... 54993 OCKHAMG-CDS TERM 55079 ... 55291 OCKHAMG-CDS

30

20

5

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 438
  - Ceres seq id 2036457

35

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 439
  - Ceres seq id 2036458
  - Location of start within SEQ ID NO 438: at 1 nt.

40

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 439: from 104 to 188

45 aa.

- (Dp) Related Amino Acid Sequences
  - Alignment No. 291
  - qi No. 133940
- Description: 40S RIBOSOMAL PROTEIN S3A (S1A) >gi|70851|pir||R3XL3A ribosomal protein S3a African clawed

260

froq >qi|65091|emb|CAA40592| (X57322) ribosomal protein Sla [Xenopus laevis]

- % Identity: 75.1
- Alignment Length: 250
- Location of Alignment in SEQ ID NO 439: from 1 to 249
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 440
- 10 - Ceres seq id 2036459
  - Location of start within SEQ ID NO 438: at 130 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Ribosomal protein S3, C-terminal domain.
    - Location within SEQ ID NO 440: from 61 to 145 aa.
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 292
      - qi No. 133940
  - Description: 40S RIBOSOMAL PROTEIN S3A (S1A) >qi|70851|pir||R3XL3A ribosomal protein S3a - African clawed froq >qi|65091|emb|CAA40592| (X57322) ribosomal protein S1a

[Xenopus laevis]

- % Identity: 75.1
- Alignment Length: 250
- Location of Alignment in SEQ ID NO 440: from 1 to
- 30 Maximum Length Sequence:

Public Genomic DNA:

gi No: 6434218

Predicted Exons:

INIT 85292 ... 85282 OCKHAMG-CDS 84903 ... 84828 OCKHAMG-CDS INTR 84730 ... 84629 OCKHAMG-CDS INTR INTR 84524 ... 84376 OCKHAMG-CDS 84282 ... 84156 OCKHAMG-CDS INTR 84148 ... 84065 OCKHAMG-CDS TERM

40

35

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15

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25

206

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 441
  - Ceres seq id 2036585

45

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 442
  - Ceres seq id 2036586
  - Location of start within SEQ ID NO 441: at 1 nt.

261 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Ribosomal protein L24e - Location within SEQ ID NO 442: from 3 to 73 aa. 5 (Dp) Related Amino Acid Sequences - Alignment No. 293 - qi No. 6094040 - Description: 60S RIBOSOMAL PROTEIN L24 10 >qi|2961300|emb|CAA12358| (AJ225027) ribosomal protein L24 [Cicer arietinum] - % Identity: 86.6 - Alignment Length: 158 - Location of Alignment in SEQ ID NO 442: from 1 to 15 156 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 443 - Ceres seq id 2036587 20 - Location of start within SEQ ID NO 441: at 166 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences 25 - Alignment No. 294 - qi No. 6094040 - Description: 60S RIBOSOMAL PROTEIN L24 >qi|2961300|emb|CAA12358| (AJ225027) ribosomal protein L24 [Cicer arietinum] 30 - % Identity: 86.6 - Alignment Length: 158 - Location of Alignment in SEQ ID NO 443: from 1 to 101 35 Maximum Length Sequence: Public Genomic DNA: qi No: 5668776 Predicted Exons: INIT 19552 ... 19704 OCKHAMG-CDS 40 TERM 19947 ... 20279 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 444 - Ceres seq id 2039554 45 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 445

- Ceres seq id 2039555
- 50 - Location of start within SEQ ID NO 444: at 1 nt.

262

- (Dp) Related Amino Acid Sequences
  - Alignment No. 295
  - qi No. 5882720

- Description: (AC008263) Similar to gb|D86180 phosphoribosylanthranilate transferase from Pisum sativum and contains 2 PF|00168 C2 (phospholipid binding) domains. ESTs gb|H76726, gb|T45544 and gb|N96377 come from this gene.

10 [Arab...

- % Identity: 99.3
- Alignment Length: 151
- Location of Alignment in SEQ ID NO 445: from 1 to

151

15

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 446
  - Ceres seq id 2039556
  - Location of start within SEQ ID NO 444: at 136 nt.

20

25

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 296
- qi No. 5882720
- Description: (AC008263) Similar to gb|D86180 phosphoribosylanthranilate transferase from Pisum sativum and contains 2 PF:00168 C2 (phospholipid binding) domains. ESTs gb|H76726, gb|T45544 and gb|N96377 come from this gene.

30 [Arab...

- % Identity: 99.3
- Alignment Length: 151
- Location of Alignment in SEQ ID NO 446: from 1 to

106

35

Maximum Length Sequence:

Public Genomic DNA:

gi No: 4263586

Predicted Exons:

40 INIT 44660 ... 44729 OCKHAMG-CDS
INTR 44822 ... 44985 OCKHAMG-CDS
INTR 45066 ... 45099 OCKHAMG-CDS
INTR 45171 ... 45264 OCKHAMG-CDS
TERM 45352 ... 45367 OCKHAMG-CDS

45

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 447
  - Ceres seq id 2044283

50

(B) Polypeptide Sequence

263

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- Pat. Appln. SEQ ID NO 448
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- Ceres seq id 2044284
- Location of start within SEQ ID NO 447: at 1 nt.
- Location of Signal Peptide Cleavage Site within SEQ 5 ID NO 448: at 19 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
- Alignment No. 297
  - gi No. 4512613
  - Description: (AC004793) F28K20.12 [Arabidopsis

thaliana]

- % Identity: 99.2
- Alignment Length: 122
  - Location of Alignment in SEQ ID NO 448: from 1 to 122
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 449
      - Ceres seq id 2044285
      - Location of start within SEQ ID NO 447: at 13 nt.
- (C) Nomination and Annotation of Domains within
- 25 Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 298
    - gi No. 4512613
    - Description: (AC004793) F28K20.12 [Arabidopsis
- 30 thaliana]
- % Identity: 99.2
- Alignment Length: 122
- Location of Alignment in SEQ ID NO 449: from 1 to

118

35

20

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 450
  - Ceres seq id 2044286
  - Location of start within SEQ ID NO 447: at 16 nt.

40

45

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 299
  - gi No. 4512613
    - Description: (AC004793) F28K20.12 [Arabidopsis

thaliana]

- % Identity: 99.2
- Alignment Length: 122
- Location of Alignment in SEQ ID NO 450: from 1 to

PCT/US00/00466

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Maximum Length Sequence:
      related to:
     Clone IDs:
 5
           157859
     Public Genomic DNA:
           gi No: 4519187
           Predicted Exons:
              INIT
                   64212 ... 64316 OCKHAMG-CDS
              INTR 64435 ... 64599 OCKHAMG-CDS
10
              INTR 64655 ... 64789 OCKHAMG-CDS
              INTR 64869 ... 64952 OCKHAMG-CDS
              INTR 65028 ... 65143 OCKHAMG-CDS
             INTR 65245 ... 65346 OCKHAMG-CDS INTR 65417 ... 65530 OCKHAMG-CDS
15
             INTR
                    65621 ... 65677 OCKHAMG-CDS
             INTR 65790 ... 65856 OCKHAMG-CDS
             INTR 65929 ... 65982 OCKHAMG-CDS
             INTR 66089 ... 66163 OCKHAMG-CDS
20
             INTR 66558 ... 66593 OCKHAMG-CDS
              INTR 66699 ... 66811 OCKHAMG-CDS
             TERM 68086 ... 68098 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 451
25
          - Ceres seq id 2048114
          - Alternative transcription start site(s) located in SEQ
     ID NO 451:
          - Clone 157859 starts at 456 and ends at in cDNA.
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 452
            - Ceres seq id 2048115
            - Location of start within SEQ ID NO 451: at 1 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
35
     ID NO 452: at 34 aa.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
40
           (Dp) Related Amino Acid Sequences
               - Alignment No. 300
               - gi No. 4850384
              - Description: (AC007357) F3F19.3 [Arabidopsis
     thaliana]
45
               - % Identity: 66.2
              - Alignment Length: 201
               - Location of Alignment in SEQ ID NO 452: from 95
    to 295
50
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 453
```

265

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- Ceres seq id 2048116
```

- Location of start within SEQ ID NO 451: at 19 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO  $453\colon$  at 28 aa.

5

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:continuous}$ 
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 301
- qi No. 4850384
  - Description: (AC007357) F3F19.3 [Arabidopsis

thaliana]

- % Identity: 66.2
- Alignment Length: 201
- Location of Alignment in SEQ ID NO 453: from 89 to 289
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 454
- 20 Ceres seq id 2048117
  - Location of start within SEQ ID NO 451: at 22 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 454: at 27 aa.
- 25 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 302
    - qi No. 4850384
- Description: (AC007357) F3F19.3 [Arabidopsis

thaliana]

- % Identity: 66.2
- Alignment Length: 201
- Location of Alignment in SEQ ID NO 454: from 88

35 to 288

50

Maximum Length Sequence:

Public Genomic DNA:

qi No: 4519194

40 Predicted Exons:

INTR 15272 ... 15482 OCKHAMG-CDNA INTR 15570 ... 15615 OCKHAMG-CDNA INTR 16443 ... 16834 OCKHAMG-CDNA

45 INIT 15350 ... 15482 OCKHAMG-CDS INTR 15570 ... 15615 OCKHAMG-CDS TERM 16443 ... 16671 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 455
  - Ceres seq id 2048271

5	<ul><li>(B) Polypeptide Sequence</li><li>- Pat. Appln. SEQ ID NO 456</li><li>- Ceres seq_id 2048272</li><li>- Location of start within SEQ ID NO 455: at 79 nt.</li></ul>
10	(C) Nomination and Annotation of Domains within Predicted Polypeptics(s) - Ribosomal protein S16 - Location within SEQ ID NO 456: from 9 to 70 aa.
15	(Dp) Related Amino Acid Sequences - Alignment No. 303 - gi No. 3861401 - Description: (AJ235273) 30S RIBOSOMAL PROTEIN S16 (rpsP) [Rickettsia prowazekii]
20	- % Identity: 43.7 - Alignment Length: 103 - Location of Alignment in SEQ ID NO 456: from 1 to
25	<ul> <li>(B) Polypeptide Sequence</li> <li>- Pat. Appln. SEQ ID NO 457</li> <li>- Ceres seq_id 2048273</li> <li>- Location of start within SEQ ID NO 455: at 139 nt.</li> </ul>
30	<ul><li>(C) Nomination and Annotation of Domains within</li><li>Predicted Polypeptide(s)</li><li>Ribosomal protein S16</li><li>Location within SEQ ID NO 457: from 1 to 50 aa.</li></ul>
35	(Dp) Related Amino Acid Sequences - Alignment No. 304 - gi No. 3861401 - Description: (AJ235273) 30S RIBOSOMAL PROTEIN S16 (rpsP) [Rickettsia prowazekii]
40	<ul> <li>- % Identity: 43.7</li> <li>- Alignment Length: 103</li> <li>- Location of Alignment in SEQ ID NO 457: from 1 to</li> <li>82</li> </ul>
45	<ul> <li>(B) Polypeptide Sequence</li> <li>- Pat. Appln. SEQ ID NO 458</li> <li>- Ceres seq_id 2048274</li> <li>- Location of start within SEQ ID NO 455: at 232 nt.</li> </ul>
50	<ul><li>(C) Nomination and Annotation of Domains within</li><li>Predicted Polypeptide(s)</li><li>(Dp) Related Amino Acid Sequences</li><li>Alignment No. 305</li></ul>

267

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- qi No. 3861401
```

- Description: (AJ235273) 30S RIBOSOMAL PROTEIN S16

(rpsP) [Rickettsia prowazekii]

- % Identity: 43.7
- Alignment Length: 103
- Location of Alignment in SEQ ID NO 458: from 1 to

51

5

15

20

25

35

Maximum Length Sequence:

10 Public Genomic DNA:

gi No: 4519195

Predicted Exons:

INIT 48116 ... 48096 OCKHAMG-CDS
INTR 48033 ... 47989 OCKHAMG-CDS
INTR 47604 ... 47531 OCKHAMG-CDS
INTR 47435 ... 47386 OCKHAMG-CDS
TERM 47286 ... 47192 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
- Pat. Appln. SEQ ID NO 459
  - Ceres seq id 2048331
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 460
    - Ceres seq id 2048332
    - Location of start within SEQ ID NO 459: at 1 nt.
- (C) Nomination and Annotation of Domains within 30 Predicted Polypeptide(s)
  - Domain of unknown function
  - Location within SEQ ID NO 460: from 8 to 92 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 306
    - gi No. 5032215
    - Description: ref|NP\_005827.1|pUK114|

translational inhibitor protein p14.5

>gi|1717975|sp|P52758|UK14 HUMAN 14.5 KD TRANSLATIONAL

40 INHIBITOR PROTEIN (P14.5) (UK114 ANTIGEN HOMOLOG)

>qi|1177435|emb|CAA64670| sapiens}

- % Identity: 58.8
- Alignment Length: 85
- Location of Alignment in SEQ ID NO 460: from 8 to

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 461
  - Ceres seq id 2048333
- 50 Location of start within SEQ ID NO 459: at 79 nt.

268

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Domain of unknown function
               - Location within SEQ ID NO 461: from 1 to 66 aa.
 5
           (Dp) Related Amino Acid Sequences
               - Alignment No. 307
               - gi No. 5032215
               - Description: ref|NP 005827.1|pUK114|
10
     translational inhibitor protein p14.5
     >gi|1717975|sp|P52758|UK14 HUMAN 14.5 KD TRANSLATIONAL
     INHIBITOR PROTEIN (P14.5) (UK114 ANTIGEN HOMOLOG)
>gi|1177435|emb|CAA64670| sapiens]
               - % Identity: 58.8
15
               - Alignment Length: 85
               - Location of Alignment in SEQ ID NO 461: from 1 to
     66
        (B) Polypeptide Sequence
20
            - Pat. Appln. SEQ ID NO 462
            - Ceres seq id 2048334
            - Location of start within SEQ ID NO 459: at 139 nt.
           (C) Nomination and Annotation of Domains within
25
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 308
               - qi No. 5032215
               - Description: ref|NP 005827.1|pUK114|
     translational inhibitor protein p14.5
30
     >gi|1717975|sp|P52758|UK14 HUMAN 14.5 KD TRANSLATIONAL
     INHIBITOR PROTEIN (P14.5) (UK114 ANTIGEN HOMOLOG)
     >gi|1177435|emb|CAA64670| sapiens]
               - % Identity:
                              58.8
35
               - Alignment Length: 85
               - Location of Alignment in SEQ ID NO 462: from 1 to
     46
     Maximum Length Sequence:
40
     Public Genomic DNA:
           qi No: 4589412
           Predicted Exons:
                     37105 ... 37091 OCKHAMG-CDS
              INIT
              INTR
                     36413 ... 36339 OCKHAMG-CDS
              INTR 36224 ... 36176 OCKHAMG-CDS
45
              INTR 35977 ... 35698 OCKHAMG-CDS
              INTR 35603 ... 35471 OCKHAMG-CDS
              TERM 35193 ... 35119
                                        OCKHAMG-CDS
```

(Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 463

269

- Ceres seq id 2048466

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(B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 464
- Ceres seq id 2048467
- Location of start within SEQ ID NO 463: at 1 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 309
    - gi No. 3688432
    - Description: (AJ011705) 40S ribosomal protein S10
- 15 [Lumbricus rubellus]
  - % Identity: 50.6
  - Alignment Length: 164
  - Location of Alignment in SEQ ID NO 464: from 30

to 193

20

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 465
  - Ceres seq id 2048468
  - Location of start within SEQ ID NO 463: at 4 nt.

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- - (Dp) Related Amino Acid Sequences
    - Alignment No. 310
    - gi No. 3688432
- Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]
  - % Identity: 50.6
  - Alignment Length: 164
- Location of Alignment in SEQ ID NO 465: from 29

to 192

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 466
  - Ceres seq id 2048469
  - Location of start within SEQ ID NO 463: at 88 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
- 45 (Dp) Related Amino Acid Sequences
  - Alignment No. 311
  - gi No. 3688432
  - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus]
  - % Identity: 50.6
    - Alignment Length: 164

270

- Location of Alignment in SEQ ID NO 466: from 1 to 164

Maximum Length Sequence:

5 Public Genomic DNA:

gi No: 4914400

Predicted Exons:

INTR 97410 ... 97833 OCKHAMG-CDNA

10 SINGLE 97525 ... 97797 OCKHAMG-CDS

gi No: 4938473
Predicted Exons:

SINGLE 9769 ... 10041 GENBANK

15 INTR 9654 ... 10078 OCKHAMG-CDNA INTR 10721 ... 10880 OCKHAMG-CDNA

SINGLE 9769 ... 10041 OCKHAMG-CDS

- 20 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 467
  - Ceres seq id 2050485
- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 468
  - Ceres seq id 2050486
  - Location of start within SEQ ID NO 467: at 2 nt.
- 30 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 312
    - qi No. 2493089
    - Description: ATP SYNTHASE F CHAIN, MITOCHONDRIAL
    - % Identity: 29.7
    - Alignment Length: 77
  - Location of Alignment in SEQ ID NO 468: from 52

to 125

35

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 469
  - Ceres seq id 2050487
  - Location of start within SEQ ID NO 467: at 116 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 313
    - qi No. 2493089
    - Description: ATP SYNTHASE F CHAIN, MITOCHONDRIAL

27Ĩ - % Identity: 29.7 - Alignment Length: 77 - Location of Alignment in SEQ ID NO 469: from 14 to 87 5 Maximum Length Sequence: Public Genomic DNA: qi No: 4454022 Predicted Exons: 55210 ... 55024 GENBANK 10 INIT 54477 ... 54209 TERM GENBANK INTR 55270 ... 55024 OCKHAMG-CDNA INTR 54477 ... 54299 OCKHAMG-CDNA 15 INIT 55210 ... 55024 OCKHAMG-CDS 54477 ... 54209 OCKHAMG-CDS TERM (Ac) cDNA Polynucleotide Sequence 20 - Pat. Appln. SEQ ID NO 470 - Ceres seq id 2050708 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 471 25 - Ceres seq id 2050709 - Location of start within SEQ ID NO 470: at 61 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) 30 (Dp) Related Amino Acid Sequences - Alignment No. 314 - qi No. 2129641 - Description: major latex protein type 1 -35 Arabidopsis thaliana >gi|1107493|emb|CAA63026| (X91960) major latex protein type1 [Arabidopsis thaliana] - % Identity: 71.3 - Alignment Length: 150 - Location of Alignment in SEQ ID NO 471: from 1 to 40 150 Maximum Length Sequence: Public Genomic DNA: gi No: 4914454 45 Predicted Exons: 42446 ... 41456 OCKHAMG-CDS INTR 40358 ... 40280 OCKHAMG-CDS INTR INTR 39551 ... 39481 OCKHAMG-CDS 39415 ... 39322 OCKHAMG-CDS INTR

INTR 39280 ... 39268 OCKHAMG-CDS INTR 38937 ... 38658 OCKHAMG-CDS

5

10

15

20

25

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45

to 560

to 520

272 INTR 38575 ... 38455 OCKHAMG-CDS TERM 38300 ... 38220 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 472 - Ceres seq id 2050901 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 473 - Ceres seq id 2050902 - Location of start within SEQ ID NO 472: at 3 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 315 - qi No. 3688432 - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus] - % Identity: 53.1 - Alignment Length: 147 - Location of Alignment in SEQ ID NO 473: from 416 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 474 - Ceres seq id 2050903 - Location of start within SEQ ID NO 472: at 123 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 316 - qi No. 3688432 - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus] - % Identity: 53.1 - Alignment Length: 147 - Location of Alignment in SEQ ID NO 474: from 376 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 475 - Ceres seq id 2050904 - Location of start within SEQ ID NO 472: at 678 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 475: at 39 aa.

50 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)

273 (Dp) Related Amino Acid Sequences - Alignment No. 317 - qi No. 3688432 - Description: (AJ011705) 40S ribosomal protein S10 [Lumbricus rubellus] 5 - % Identity: 53.1 - Alignment Length: 147 - Location of Alignment in SEQ ID NO 475: from 191 to 335 10 Maximum Length Sequence: Public Genomic DNA: qi No: 4725940 Predicted Exons: 15 SINGLE 96089 ... 95736 GENBANK INTR 96156 ... 95622 OCKHAMG-CDNA SINGLE 96083 ... 95736 OCKHAMG-CDS gi No: 5823567 20 Predicted Exons: SINGLE 96089 ... 95736 GENBANK INTR 96156 ... 95622 OCKHAMG-CDNA 25 SINGLE 96083 ... 95736 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 476 30 - Ceres seq id 2051325 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 477 35 - Ceres seq id 2051326 - Location of start within SEQ ID NO 476: at 68 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 477: at 30 aa. (C) Nomination and Annotation of Domains within 40 Predicted Polypeptide(s) - Plant lipid transfer protein family - Location within SEQ ID NO 477: from 34 to 115 aa. 45 (Dp) Related Amino Acid Sequences - Alignment No. 318 - qi No. 4725952 - Description: (AL049730) AIR1A-like protein [Arabidopsis thaliana]

- % Identity: 100

- Alignment Length: 117

274 - Location of Alignment in SEQ ID NO 477: from 1 to 117 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 478 - Ceres seq id 2051327 - Location of start within SEQ ID NO 476: at 74 nt. - Location of Signal Peptide Cleavage Site within SEO ID NO 478: at 28 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Plant lipid transfer protein family - Location within SEQ ID NO 478: from 32 to 113 aa. (Dp) Related Amino Acid Sequences - Alignment No. 319 - qi No. 4725952 - Description: (AL049730) AIR1A-like protein [Arabidopsis thaliana] - % Identity: 100 - Alignment Length: 117 - Location of Alignment in SEQ ID NO 478: from 1 to 115 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 479 - Ceres seq id 2051328 - Location of start within SEQ ID NO 476: at 288 nt. - Location of Signal Peptide Cleavage Site within SEO ID NO 479: at 61 aa. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences Maximum Length Sequence: Public Genomic DNA: qi No: 4490717 Predicted Exons: 87492 ... INIT 87378 GENBANK 87034 ... 86928 GENBANK INTR INTR 86683 ... 86641 **GENBANK** TERM 86445 ... 86402 **GENBANK** 87555 ... 87378 OCKHAMG-CDNA INTR 87034 ... 86928 OCKHAMG-CDNA INTR

INTR 86683 ... 86641 OCKHAMG-CDNA

INIT 87492 ... 87378 OCKHAMG-CDS

INTR

86445 ... 86257 OCKHAMG-CDNA

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275
              INTR 87034 ...
                                86928 OCKHAMG-CDS
              INTR 86683 ...
                                86641 OCKHAMG-CDS
              TERM
                     86445 ... 86402 OCKHAMG-CDS
5
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 480
          - Ceres seq id 2051633
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 481
            - Ceres seq id 2051634
            - Location of start within SEQ ID NO 480: at 1 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 320
               - qi No. 1083282
20
               - Description: cytochrome-c oxidase (EC 1.9.3.1) -
     mouse >gi|567766 (L06465) cytochrome c oxidase [Mus musculus]
     >gi|1094404|prf||2106151A cytochrome c oxidase:SUBUNIT=VIa
     [Mus musculus]
               - % Identity: 41.4
25
               - Alignment Length: 106
               - Location of Alignment in SEQ ID NO 481: from 22
     to 120
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 482
30
            - Ceres seq id 2051635
            - Location of start within SEQ ID NO 480: at 64 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
35
           (Dp) Related Amino Acid Sequences
               - Alignment No. 321
               - qi No. 1083282
               - Description: cytochrome-c oxidase (EC 1.9.3.1) -
     mouse >qi|567766 (L06465) cytochrome c oxidase [Mus musculus]
40
     >gi|1094404|prf||2106151A cytochrome c oxidase:SUBUNIT=VIa
     [Mus musculus]
               - % Identity: 41.4
               - Alignment Length: 106
45
               - Location of Alignment in SEQ ID NO 482: from 1 to
     99
     Maximum Length Sequence:
     Public Genomic DNA:
           gi No: 4584531
50
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Predicted Exons:

```
276
                                56641 OCKHAMG-CDNA
              INTR 56873 ...
                    56627 ... 56451
              INTR
                                        OCKHAMG-CDNA
              SINGLE 56812 ... 56645
                                        OCKHAMG-CDS
5
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 483
          - Ceres seq id 2051906
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 484
            - Ceres seq id 2051907
            - Location of start within SEQ ID NO 483: at 2 nt.
15
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
               - Alignment No. 322
20
               - qi No. 4850407
               - Description: (AC007357) EST gb|T21221 comes from
     this gene. [Arabidopsis thaliana]
               - % Identity: 57.1
               - Alignment Length: 35
25
               - Location of Alignment in SEQ ID NO 484: from 37
     to 70
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 485
30
            - Ceres seq id 2051908
            - Location of start within SEQ ID NO 483: at 62 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
35
               - Alignment No. 323
               - qi No. 4850407
               - Description: (AC007357) EST gb|T21221 comes from
     this gene. [Arabidopsis thaliana]
40
               - % Identity: 57.1
               - Alignment Length: 35
               - Location of Alignment in SEQ ID NO 485: from 17
     to 50
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 486
            - Ceres seq id 2051909
            - Location of start within SEQ ID NO 483: at 74 nt.
50
           (C) Nomination and Annotation of Domains within
```

Predicted Polypeptide(s)

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277
           (Dp) Related Amino Acid Sequences
               - Alignment No. 324
               - qi No. 4850407
               - Description: (AC007357) EST gb|T21221 comes from
5
    this gene. [Arabidopsis thaliana]
               - % Identity: 57.1
               - Alignment Length: 35
               - Location of Alignment in SEQ ID NO 486: from 13
    to 46
10
    Maximum Length Sequence:
     Public Genomic DNA:
           qi No: 4584841
           Predicted Exons:
15
              INTR
                    75983 ...
                                75751 OCKHAMG-CDNA
                    75412 ... 74936 OCKHAMG-CDNA
              INTR
                    75992 ... 75751 OCKHAMG-CDNA
              INTR
                               75090 OCKHAMG-CDNA
                    75412 ...
              INTR
20
                     75937 ... 75751 OCKHAMG-CDS
              INIT
              TERM
                     75412 ... 75132
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
25
          - Pat. Appln. SEQ ID NO 487
          - Ceres seq id 2052403
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 488
30
            - Ceres seq id 2052404
            - Location of start within SEQ ID NO 487: at 56 nt.
           (C) Nomination and Annotation of Domains within
35
     Predicted Polypeptide(s)
               - Pathogenesis-related protein Bet v I family
               - Location within SEQ ID NO 488: from 5 to 155 aa.
           (Dp) Related Amino Acid Sequences
40
               - Alignment No. 325
               - qi No. 1321731
               - Description: (Z72439) major allergen Cor a 1
     [Corylus avellana]
               - % Identity:
                              36.3
45
               - Alignment Length: 159
               - Location of Alignment in SEQ ID NO 488: from 5 to
     155
     Maximum Length Sequence:
```

50

Public Genomic DNA:

gi No: 4699904

278 Predicted Exons: INTR 82618 ... 82845 OCKHAMG-CDNA INTR 83226 ... 83301 OCKHAMG-CDNA INTR 83389 ... 83561 OCKHAMG-CDNA INTR 84124 ... 84385 OCKHAMG-CDNA 5 INIT 82730 ... 82845 OCKHAMG-CDS INTR 83226 ... 83301 OCKHAMG-CDS INTR 83389 ... 83561 OCKHAMG-CDS 84124 ... 84199 OCKHAMG-CDS 10 TERM (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 489 - Ceres seq id 2053545 15 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 490 - Ceres seq id 2053546 20 - Location of start within SEQ ID NO 489: at 113 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 490: at 36 aa. (C) Nomination and Annotation of Domains within 25 Predicted Polypeptide(s) - MAPEG family - Location within SEQ ID NO 490: from 59 to 136 aa. (Dp) Related Amino Acid Sequences - Alignment No. 326 30 - qi No. 4758714 - Description: ref|NP 004519.1|pMGST3| microsomal glutathione S-transferase 3 >gi|2583081 (AF026977) microsomal

- glutathione S-transferase 3 [Homo sapiens]
  - % Identity: 41.9
    - Alignment Length: 131
- Location of Alignment in SEQ ID NO 490: from 8 to 136
- 40 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 491
- Ceres seg id 2053547
- Location of start within SEQ ID NO 489: at 197 nt.
- (C) Nomination and Annotation of Domains within 45 Predicted Polypeptide(s)
  - MAPEG family
  - Location within SEQ ID NO 491: from 31 to 108 aa.
- 50 (Dp) Related Amino Acid Sequences - Alignment No. 327

279

- qi No. 4758714

- Description: ref|NP\_004519.1|pMGST3| microsomal glutathione S-transferase 3 >gi|2583081 (AF026977) microsomal glutathione S-transferase 3 [Homo sapiens]

- % Identity: 41.9

- Alignment Length: 131

- Location of Alignment in SEQ ID NO 491: from 1 to 108

- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 492
  - Ceres seq id 2053548
  - Location of start within SEQ ID NO 489: at 329 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 15 ID NO 492: at 22 aa.

5

- (C) Nomination and Annotation of Domains within  $Predicted\ Polypeptide(s)$ 
  - MAPEG family
- Location within SEQ ID NO 492: from 1 to 64 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 328
    - qi No. 4758714
- Description: ref[NP\_004519.1|pMGST3| microsomal glutathione S-transferase 3 >gi|2583081 (AF026977) microsomal glutathione S-transferase 3 [Homo sapiens]
  - % Identity: 41.9
  - Alignment Length: 131
- Location of Alignment in SEQ ID NO 492: from 1 to

Maximum Length Sequence:

related to:

35 Clone IDs:

2004

31675

266907

Public Genomic DNA:

40 gi No: 4887737

Predicted Exons:

	INTR		8579	OCKHAMG-CDNA
45	INTR	7913	 8587	OCKHAMG-CDNA
	INTR	7911	 8579	OCKHAMG-CDNA
	INTR	7909	 8579	OCKHAMG-CDNA
50	INTR	7913	 8578	OCKHAMG-CDNA

					280	
		INTR	7913	• • •	8509	OCKHAMG-CDNA
_	SINGLE No: 503 edicted	19264		8425	OCKHAMG-CDS	
	SINGLE			4610	GENBANK	
		INTR	4092		4764	OCKHAMG-CDNA
10		INTR	4098		4772	OCKHAMG-CDNA
		INTR	4096		4764	OCKHAMG-CDNA
15		INTR	4094		4764	OCKHAMG-CDNA
10		INTR	4098		4763	OCKHAMG-CDNA
		INTR	4098		4694	OCKHAMG-CDNA
20 SINGLE 4143 4610 OCKHAMG-CDS  (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 493 - Ceres seq_id 2053884						
- Alternative transcription start site(s) located in SEQ 25 ID NO 493:						
30	<ul><li>Clone 31675 starts at 3 and ends at 607 in cDNA.</li><li>Clone 266907 starts at 4 and ends at 677 in cDNA.</li></ul>					
(B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 494 - Ceres seq_id 2053885 - Location of start within SEQ ID NO 493: at 1 nt.						
<ul><li>(C) Nomination and Annotation of Domains within</li><li>Predicted Polypeptide(s)</li><li>(Dp) Related Amino Acid Sequences</li></ul>						
40						
45	159		_			in SEQ ID NO 494: from 2 to
	(B) P	olypept			e D NO 405	

- Pat. Appln. SEQ ID NO 495 Ceres seq\_id 2053886

50

- Location of start within SEQ ID NO 493: at 52 nt.

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(C) Nomination and Annotation of Domains within
    Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
5
              - Alignment No. 330
              - qi No. 2498177
              - Description: BASP1 PROTEIN
              - % Identity:
                             28
            - Alignment Length: 164
              - Location of Alignment in SEQ ID NO 495: from 1 to
10
    142
        (B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 496
           - Ceres seq id 2053887
15
           - Location of start within SEQ ID NO 493: at 172 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
20
           (Dp) Related Amino Acid Sequences
               - Alignment No. 331
              - qi No. 2498177
              - Description: BASP1 PROTEIN
               - % Identity:
                             28
25
              - Alignment Length: 164
              - Location of Alignment in SEQ ID NO 496: from 1 to
     102
    Maximum Length Sequence:
30
    Public Genomic DNA:
          gi No: 4887738
          Predicted Exons:
              INTR 45004 ... 44725
                                       OCKHAMG-CDNA
              INTR 44038 ... 43918
                                       OCKHAMG-CDNA
35
              INTR 43839 ... 43444
                                       OCKHAMG-CDNA
              INIT 44942 ... 44725 OCKHAMG-CDS
                    44038 ... 43918
                                       OCKHAMG-CDS
              INTR
                    43839 ... 43564
              TERM
                                       OCKHAMG-CDS
40
          gi No: 5103850
           Predicted Exons:
              INIT
                  44944 ... 44727
                                       GENBANK
                    44040 ... 43920
              INTR
                                       GENBANK
                    43841 ... 43566
              TERM
                                       GENBANK
45
              INTR 45006 ... 44727 OCKHAMG-CDNA
              INTR 44040 ... 43920
                                      OCKHAMG-CDNA
                    43841 ... 43446
              INTR
                                       OCKHAMG-CDNA
50
                    44944 ... 44727 OCKHAMG-CDS
              INIT
              INTR
                    44040 ... 43920
                                      OCKHAMG-CDS
```

282

TERM 43841 ... 43566 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 497
- Ceres seq id 2053908
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 498
- Ceres seg id 2053909
  - Location of start within SEQ ID NO 497: at 63 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 332
    - qi No. 2245000
    - Description: (Z97341) LET1 like protein

[Arabidopsis thaliana]

- 20 % Identity: 31.4
  - Alignment Length: 156
  - Location of Alignment in SEQ ID NO 498: from 17

to 164

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- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 499
  - Ceres seq id 2053910
  - Location of start within SEQ ID NO 497: at 168 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 30 ID NO 499: at 23 aa.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
    - Alignment No. 333
      - gi No. 2245000
      - Description: (Z97341) LET1 like protein

[Arabidopsis thaliana]

- % Identity: 31.4
- Alignment Length: 156
- Location of Alignment in SEQ ID NO 499: from 1 to 129
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 500
    - Ceres seq id 2053911
    - Location of start within SEQ ID NO 497: at 216 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences

- Description: (Z97341) LET1 like protein

283
- Alignment No. 334
- gi No. 2245000

[Arabidopsis thaliana]

- % Identity: 31.4

- Alignment Length: 156

- Location of Alignment in SEQ ID NO 500: from 1 to

113

5

10 Maximum Length Sequence:

Public Genomic DNA:

gi No: 4335711

TERM

Predicted Exons:

INTR 46239 ... 46470 OCKHAMG-CDNA
INTR 46624 ... 46739 OCKHAMG-CDNA
INTR 47125 ... 47428 OCKHAMG-CDNA
INIT 46285 ... 46470 OCKHAMG-CDS
INTR 46624 ... 46739 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 501
- Ceres seq\_id 2056123

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- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 502
  - Ceres seq id 2056124
- Location of start within SEQ ID NO 501: at 47 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Zinc finger, C3HC4 type (RING finger)

47125 ... 47413 OCKHAMG-CDS

- Location within SEQ ID NO 502: from 150 to 190 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 335
    - qi No. 3790593
  - Description: (AF079185) RING-H2 finger protein

RHY1a [Arabidopsis thaliana]

- % Identity: 55.1
- Alignment Length: 49
- Location of Alignment in SEQ ID NO 502: from 146 to 194
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 503
    - Ceres seq id 2056125
    - Location of start within SEQ ID NO 501: at 56 nt.

```
(C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
               - Zinc finger, C3HC4 type (RING finger)
5
               - Location within SEQ ID NO 503: from 147 to 187
     aa.
           (Dp) Related Amino Acid Sequences
               - Alignment No. 336
10
               - qi No. 3790593
               - Description: (AF079185) RING-H2 finger protein
     RHYla [Arabidopsis thaliana]
               - % Identity: 55.1
             - Alignment Length: 49
15
               - Location of Alignment in SEQ ID NO 503: from 143
     to 191
    Maximum Length Sequence:
     Public Genomic DNA:
20
           gi No: 4432829
           Predicted Exons:
              INTR 51185 ...
                               51056 OCKHAMG-CDNA
              INTR 50590 ...
                               50253 OCKHAMG-CDNA
25
              INIT
                     56176 ... 55097 OCKHAMG-CDS
              INTR 55014 ... 54161 OCKHAMG-CDS
              INTR 54076 ... 54011 OCKHAMG-CDS INTR 51530 ... 51382 OCKHAMG-CDS
                     51187 ... 51056 OCKHAMG-CDS
              INTR
30
              TERM
                     50590 ... 50418 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 504
          - Ceres seq id 2056245
35
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 505
            - Ceres seq id 2056246
40
            - Location of start within SEQ ID NO 504: at 1 nt.
           (C) Nomination and Annotation of Domains within
     Predicted Polypeptide(s)
           (Dp) Related Amino Acid Sequences
45
               - Alignment No. 337
               - qi No. 4938484
               - Description: (AL078464) transcription factor-like
     protein [Arabidopsis thaliana]
               - % Identity: 56.2
50
               - Alignment Length: 661
```

285

- Location of Alignment in SEQ ID NO 505: from 4 to 644

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 506
  - Ceres seq id 2056247
  - Location of start within SEQ ID NO 504: at 55 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 338
    - qi No. 4938484
    - Description: (AL078464) transcription factor-like
- 15 protein [Arabidopsis thaliana]
  - % Identity: 56.2
  - Alignment Length: 661
  - Location of Alignment in SEQ ID NO 506: from 1 to

626 20

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 507
  - Ceres seq id 2056248
  - Location of start within SEQ ID NO 504: at 751 nt.

25

30

- - (Dp) Related Amino Acid Sequences
    - Alignment No. 339
  - gi No. 4938484
- Description: (AL078464) transcription factor-like protein [Arabidopsis thaliana]
  - % Identity: 56.2
  - Alignment Length: 661
- Location of Alignment in SEQ ID NO 507: from 1 to 394

Maximum Length Sequence:
 related to:

40 Clone IDs:

: 103.

213322

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 508
  - Ceres seq id 1941142

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 509
  - Ceres seq id 1941143
- Location of start within SEQ ID NO 508: at 119 nt.

286

(C) Nomination and Annotation of Domains within Predicted Polypeptide(s) - Ribosomal protein L24e - Location within SEO ID NO 509: from 3 to 73 aa.

- (Dp) Related Amino Acid Sequences
  - Alignment No. 340
  - qi No. 1710521
  - Description: 60S RIBOSOMAL PROTEIN L24
- >qi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein 10 [Hordeum vulgare]
  - % Identity: 94.4
  - Alignment Length: 162
  - Location of Alignment in SEQ ID NO 509: from 1 to
- 15 161

25

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 510
  - Ceres seq id 1941144
- 20 - Location of start within SEQ ID NO 508: at 284 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 341
      - gi No. 1710521
      - Description: 60S RIBOSOMAL PROTEIN L24
  - >qi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein [Hordeum vulgare]
- % Identity: 94.4 30
  - Alignment Length: 162
  - Location of Alignment in SEQ ID NO 510: from 1 to

106

35 Maximum Length Sequence:

related to:

Clone IDs:

241379

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 511
  - Ceres seq id 1942975
- Alternative transcription start site(s) located in SEQ ID NO 511:
  - 29,30,31,32,33,36

45

40

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 512
  - Ceres seg id 1942976
  - Location of start within SEQ ID NO 511: at 1 nt.

287

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 512: from 149 to 233

5 aa.

15

- (Dp) Related Amino Acid Sequences
  - Alignment No. 342
  - gi No. 133940
- Description: 40S RIBOSOMAL PROTEIN S3A (S1A) >gi|70851|pir||R3XL3A ribosomal protein S3a African clawed frog >gi|65091|emb|CAA40592| (X57322) ribosomal protein S1a [Xenopus laevis]
  - % Identity: 82.5
  - Alignment Length: 223
  - Location of Alignment in SEQ ID NO 512: from 46 to 268  $\,$ 
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 513
      - Ceres seq id 1942977
      - Location of start within SEQ ID NO 511: at 136 nt.
- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 513: from 104 to 188 aa.
- 30 (Dp) Related Amino Acid Sequences
  - Alignment No. 343
  - qi No. 133940
  - Description: 40S RIBOSOMAL PROTEIN S3A (S1A)
- >gi|70851|pir||R3XL3A ribosomal protein S3a African clawed 35 frog >gi|65091|emb|CAA40592| (X57322) ribosomal protein S1a [Xenopus laevis]
  - % Identity: 82.5
  - Alignment Length: 223
  - Location of Alignment in SEQ ID NO 513: from 1 to
- 40 223
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 514
    - Ceres seq id 1942978
- Location of start within SEQ ID NO 511: at 205 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Ribosomal protein S3, C-terminal domain.
- Location within SEQ ID NO 514: from 81 to 165 aa.

288

- (Dp) Related Amino Acid Sequences
  - Alignment No. 344
  - qi No. 133940
  - Description: 40S RIBOSOMAL PROTEIN S3A (S1A)
- 5 >gi|70851|pir||R3XL3A ribosomal protein S3a African clawed frog >gi|65091|emb|CAA40592| (X57322) ribosomal protein S1a [Xenopus laevis]
  - % Identity: 82.5
  - Alignment Length: 223
- Location of Alignment in SEQ ID NO 514: from 1 to

200

Maximum Length Sequence:

related to:

15 Clone IDs:

289536

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 515
  - Ceres seq id 1944349

20

35

50

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 516
  - Ceres seq id 1944350
- 25 Location of start within SEQ ID NO 515: at 423 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)  $\label{eq:condition}$ 
    - Ribosomal protein L24e
- Location within SEQ ID NO 516: from 3 to 73 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 345
    - gi No. 1710521
  - Description: 60S RIBOSOMAL PROTEIN L24

>gi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein
[Hordeum vulgare]

- % Identity: 94.4
- Alignment Length: 162
- Location of Alignment in SEQ ID NO 516: from 1 to 161
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 517
- 45 Ceres seq id 1944351
  - Location of start within SEQ ID NO 515: at 588 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - (Dp) Related Amino Acid Sequences
      - Alignment No. 346

289

```
- qi No. 1710521
```

- Description: 60S RIBOSOMAL PROTEIN L24

>gi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein
[Hordeum vulgare]

- % Identity: 94.4
- Alignment Length: 162
- Location of Alignment in SEQ ID NO 517: from 1 to 106

10 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 518
- Ceres seq id 1944352
- Location of start within SEQ ID NO 515: at 736 nt.
- 15 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 347
    - gi No. 1362587
- Description: spermatid-specific protein T2

precursor - longfin squid

- % Identity: 55.8
- Alignment Length: 52
- Location of Alignment in SEQ ID NO 518: from 1 to

25 46

40

45

50

5

Maximum Length Sequence:

related to:

Clone IDs:

30 291258

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 519
- Ceres seq id 1964011
- Alternative transcription start site(s) located in SEQ 35 ID NO 519:

4,34,37,38,39,40,41

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 520
- Ceres seq id 1964012
  - Location of start within SEQ ID NO 519: at 3 nt.
  - (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
    - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 520: from 150 to 234 aa.

(Dp) Related Amino Acid Sequences

- Alignment No. 348
- gi No. 1173253

290

- Description: 40S RIBOSOMAL PROTEIN S3 >gi|543317|pir||S41170 ribosomal protein S3 - mouse >gi|57728|emb|CAA35916| (X51536) ribosomal protein S3 (AA 1-243) [Rattus rattus] >gi|439522|emb|CAA54167| (X76772) ribosomal protein S3 [Mus musculus] - % Identity: 81.1 - Alignment Length: 227

- Location of Alignment in SEQ ID NO 520: from 47

to 273

(B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 521
- Ceres seq id 1964013
- Location of start within SEQ ID NO 519: at 141 nt.

15

45

10

5

- (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 521: from 104 to 188

20 aa.

- (Dp) Related Amino Acid Sequences
  - Alignment No. 349
  - qi No. 1173253
- 25 - Description: 40S RIBOSOMAL PROTEIN S3 >gi|543317|pir||S41170 ribosomal protein S3 - mouse >gi|57728|emb|CAA35916| (X51536) ribosomal protein S3 (AA 1-243) [Rattus lattus] >qi|439522|emb|CAA54167| (X76772) ribosomal protein S3 [Mus musculus] 30

- % Identity: 81.1

- Alignment Length: 227
- Location of Alignment in SEQ ID NO 521: from 1 to 227

35 (B) Polypeptide Sequence

- Pat. Appln. SEQ ID NO 522
- Ceres seg id 1964014
- Location of start within SEQ ID NO 519: at 210 nt.
- 40 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - Ribosomal protein S3, C-terminal domain.
  - Location within SEQ ID NO 522: from 81 to 165 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 350
    - qi No. 1173253
    - Description: 40S RIBOSOMAL PROTEIN S3
  - >gi|543317|pir||S41170 ribosomal protein S3 mouse
- 50 >gi|57728|emb|CAA35916| (X51536) ribosomal protein S3 (AA 1-

291 243) [Rattus rattus] >qi|439522|emb|CAA54167| (X76772) ribosomal protein S3 [Mus musculus] - % Identity: 81.1 - Alignment Length: 227 5 - Location of Alignment in SEQ ID NO 522: from 1 to 204 Maximum Length Sequence: related to: 10 Clone IDs: 207986 224937 227089 229580 15 241662 275880 278084 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 523 20 - Ceres seq id 1983854 - Alternative transcription start site(s) located in SEQ ID NO 523: -1,2,6,30,31,32,33,34,36 - Clone 207986 starts at 31 and ends at 810 in cDNA. 25 - Clone 224937 starts at 31 and ends at 810 in cDNA. - Clone 227089 starts at 33 and ends at 804 in cDNA. - Clone 229580 starts at 31 and ends at 821 in cDNA. - Clone 241662 starts at 34 and ends a 815 in cDNA. - Clone 275880 starts at 1 and ends at 804 in cDNA. - Clone 278084 starts at 31 and ends at 815 in cDNA. 30 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 524 - Ceres seq id 1983855 35 - Location of start within SEQ ID NO 523: at 315 nt. (C) Nomination and Annotation of Domains within Predicted Polypeptide(s) (Dp) Related Amino Acid Sequences - Alignment No. 351 40 - qi No. 1710521 - Description: 60S RIBOSOMAL PROTEIN L24 >qi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein [Hordeum vulgare] - % Identity: 94 45 - Alignment Length: 150 - Location of Alignment in SEQ ID NO 524: from 1 to

Maximum Length Sequence: 50 related to:

292

Clone IDs:

10

15

25

30

221977

226126

293001

- 5 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 525
  - Ceres seq id 1990261
  - Alternative transcription start site(s) located in SEQ ID NO 525:

16,30,32

- ~ Clone 226126 starts at 30 and ends at 772 in cDNA.
- Clone 293001 starts at 32 and ends at 810 in cDNA.
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 526
  - Ceres seq id 2061972
  - Location of start within SEQ ID NO 525: at 144 nt.
- (C) Nomination and Annotation of Domains within 20 Predicted Polypeptide(s)
  - Ribosomal protein L24e
  - Location within SEQ ID NO 526: from 3 to 73 aa.
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 352
      - gi No. 1710521
      - Description: 60S RIBOSOM, L PROTEIN L24

>gi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein [Hordeum vulgare]

- % Identity: 94.4
  - Alignment Length: 162
  - Location of Alignment in SEQ ID NO 526: from 1 to 161
- 35 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 527
  - Ceres seq id 2061973
  - Location of start within SEQ ID NO 525: at 309 nt.
- 40 (C) Nomination and Annotation of Domains within Predicted Polypeptide(s)
  - (Dp) Related Amino Acid Sequences
    - Alignment No. 353
    - qi No. 1710521
- Description: 60S RIBOSOMAL PROTEIN L24 >gi|1154859|emb|CAA63960| (X94296) L24 ribosomal protein [Hordeum vulgare]
  - % Identity: 94.4
  - Alignment Length: 162
- Location of Alignment in SEQ ID NO 527: from 1 to 106

293

```
REF TABLE 2
     Maximum Length Sequence:
      related to:
 5
     Clone IDs:
           9458
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 1
          - Ceres seq id 1007802
10
          - Alternative transcription start site(s) located in SEQ
     ID NO 1:
            2,3,4
        (B) Polypeptide Sequence
15
            - Pat. Appln. SEQ ID NO 2
            - Ceres seq_id 1007803
            - Location of start within SEQ ID NO 1: at 3 nt.
        (B) Polypeptide Sequence
20
            - Pat. Appln. SEQ ID NO 3
            - Ceres seq id 1007804
            - Location of start within SEQ ID NO 1: at 2 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 3: at 29 aa.
25
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 4
            - Ceres seq id 1007805
            - Location of start within SEQ ID NO 1: at 29 nt.
30
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 4: at 20 aa.
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
35
                                         specific gene, plant
     specific gene.
    Maximum Length Sequence:
      related to:
40
     Clone IDs:
           8458
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 5
          - Ceres seq id 1008556
          - Alternative transcription start site(s) located in SEQ
45
     ID NO 5:
            80, 103, 134, 135, 136, 137, 138, 139, 152, 153, 157, 166, 168
        (B) Polypeptide Sequence
50
            - Pat. Appln. SEQ ID NO 6
```

- Ceres seq id 1008557

294 - Location of start within SEQ ID NO 5: at 173 nt. (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot 5 specific gene, plant specific gene. Maximum Length Sequence: related to: Clone IDs: 10 8384 (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 7 - Ceres seq id 1008628 15 - Alternative transcription start site(s) located in SEQ ID NO 7: 3,28 (B) Polypeptide Sequence 20 - Pat. Appln. SEQ ID NO 8 - Ceres seq id 1008629 - Location of start within SEQ ID NO 7: at 41 nt. (Ba) Polypeptide Activities: Arabidopsis specific 25 gene, dicot specific gene, plant specific gene. Maximum Length Sequence: 30 related to: Clone IDs: 7260 Public Genomic DNA: qi No: 5733889 35 Predicted Exons: INIT 21286 ... 21509 GENBANK INTR 21606 ... 21832 GENBANK TERM 21951 ... 21958 GENBANK INIT 21286 ... 21509 OCKHAMG-CDS 40 21606 ... 21832 OCKHAMG-CDS INTR TERM 21951 ... 21958 OCKHAMG-CDS gi No: 6041810 Predicted Exons: 34915 ... 35138 OCKHAMG-CDS 45 INIT 35235 ... 35461 OCKHAMG-CDS INTR TERM 35580 ... 35587 OCKHAMG-CDS

100255... 100478 OCKHAMG-CDS

100575... 100801 OCKHAMG-CDS

gi No: 6091711 Predicted Exons:

> INIT INTR

```
295
              TERM 100920... 100927 OCKHAMG-CDS
           qi No: 6102641
           Predicted Exons:
              INIT
                    91941 ... 92164 OCKHAMG-CDS
                     92261 ... 92487 OCKHAMG-CDS
5
              INTR
              TERM 92606 ... 92613 OCKHAMG-CDS
           qi No: 6453849
           Predicted Exons:
                    19658 ... 19435 OCKHAMG-CDS
              INIT
10
                    19338 ... 19112
              INTR
                                       OCKHAMG-CDS
                    18993 ... 18986 OCKHAMG-CDS
              TERM
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 9
          - Ceres seq id 1009376
15
          - Alternative transcription start site(s) located in SEO
     ID NO 9:
            4,7,22,48,56,77
        (B) Polypeptide Sequence
20
            - Pat. Appln. SEQ ID NO 10
            - Ceres seq id 1009377
            - Location of start within SEQ ID NO 9: at 238 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 10: at 76 aa.
25
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 11
            - Ceres seq id 1009378
            - Location of start within SEQ ID NO 9: at 334 nt.
            - Location of Signal Peptide Cleavage Site within SEO
30
    ID NO 11: at 44 aa.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 12
35
            - Ceres seq id 1009379
            - Location of start within SEQ ID NO 9: at 245 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 12: at 47 aa.
40
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
                                        specific gene, plant
     specific gene.
45
    Maximum Length Sequence:
      related to:
     Clone IDs:
           4489
50
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 13
```

```
296
          - Ceres seq id 1011128
          - Alternative transcription start site(s) located in SEQ
     ID NO 13:
            6,7
 5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 14
            - Ceres seq id 1011129
            - Location of start within SEQ ID NO 13: at 41 nt.
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 15
            - Ceres seq id 1011130
            - Location of start within SEQ ID NO 13: at 59 nt.
15
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
                                         specific gene, plant
     specific gene.
20
     Maximum Length Sequence:
      related to:
     Clone IDs:
25
           42241
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 16
          - Ceres seq id 1011718
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 17
            - Ceres seq id 1011719
            - Location of start within SEQ ID NO 16: at 3 nt.
35
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 18
            - Ceres seq id 1011720
            - Location of start within SEQ ID NO 16: at 78 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
40
     ID NO 18: at 21 aa.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 19
            - Ceres seq id 1011721
45
            - Location of start within SEQ ID NO 16: at 102 nt.
     (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
```

specific gene, plant specific gene.

297

```
Maximum Length Sequence:
      related to:
 5
     Clone IDs:
           42200
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 20
          - Ceres seq id 1011735
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 21
            - Ceres seq id 1011736
15
            - Location of start within SEQ ID NO 20: at 75 nt.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 22
            - Ceres seq id 1011737
            - Location of start within SEQ ID NO 20: at 1 nt.
20
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 22: at 37 aa.
        (B) Polypeptide Sequence
25
            - Pat. Appln. SEQ ID NO 23
            - Ceres seq id 1011738
            - Location of start within SEQ ID NO 20: at 2 nt.
        (Ba) Polypeptide Activities: Arabidopsis specific gene,
30
     dicot
                                         specific gene, plant
     specific gene.
     Maximum Length Sequence:
35
     related to:
     Clone IDs:
           42217
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 24
40
          - Ceres seq id 1011755
          - Alternative transcription start site(s) located in SEQ
     ID NO 24:
            34
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 25
            - Ceres seq id 1011756
            - Location of start within SEQ ID NO 24: at 1 nt.
50
        (B) Polypeptide Sequence
```

- Pat. Appln. SEQ ID NO 26

298

```
- Ceres seq id 1011757
```

- Location of start within SEQ ID NO 24: at 19 nt.

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

Maximum Length Sequence:

10 related to:

Clone IDs:

42034

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 27
- 15 Ceres seq id 1011832
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 28
    - Ceres seq id 1011833
- 20 Location of start within SEQ ID NO 27: at 97 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 29
    - Ceres seq id 1011834
- 25 Location of start within SEQ ID NO 27: at 100 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 30
    - Ceres seq id 1011835
- 30 Location of start within SEQ ID NO 27: at 106 nt.
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

35 specific gene.

Maximum Length Sequence:

related to:

40 Clone IDs:

41812

Public Genomic DNA:

gi No: 6434227

Predicted Exons:

45 INIT 65854 ... 65974 OCKHAMG-CDS TERM 66067 ... 66206 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 31
  - Ceres seq id 1011907

50

(B) Polypeptide Sequence

WO 00/40695

PCT/US00/00466 299 - Pat. Appln. SEQ ID NO 32 - Ceres seq id 1011908 - Location of start within SEQ ID NO 31: at 218 nt. - Location of Signal Peptide Cleavage Site within SEQ 5 ID NO 32: at 30 aa. (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 33 - Ceres seq id 1011909 10 - Location of start within SEQ ID NO 31: at 239 nt. - Location of Signal Peptide Cleavage Site within SEO ID NO 33: at 23 aa. (B) Polypeptide Sequence 15 - Pat. Appln. SEQ ID NO 34 - Ceres seq id 1011910 - Location of start within SEQ ID NO 31: at 3 nt. - Location of Signal Peptide Cleavage Site within SEQ ID NO 34: at 17 aa. 20 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene. 25 Maximum Length Sequence: related to: Clone IDs: 41828 30 Public Genomic DNA: qi No: 4159707 Predicted Exons: 15084 ... 15201 OCKHAMG-CDNA INTR INTR 15295 ... 15324 OCKHAMG-CDNA 15414 ... 15533 OCKHAMG-CDNA 35 INTR 15648 ... 15956 OCKHAMG-CDNA INTR (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 35 - Ceres seq id 1011911 40 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 36 - Ceres seq id 1011912 - Location of start within SEQ ID NO 35: at 194 nt. 45 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 37 - Ceres seq id 1011913

- Location of start within SEQ ID NO 35: at 230 nt. - Location of Signal Peptide Cleavage Site within SEQ 50 ID NO 37: at 22 aa.

WO 00/40695

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 38
            - Ceres seq id 1011914
5
            - Location of start within SEQ ID NO 35: at 2 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 38: at 28 aa.
        (Ba) Polypeptide Activities: Arabidopsis specific gene,
10
    dicot
                                        specific gene, plant
 specific gene.
15
    Maximum Length Sequence:
     related to:
    Clone IDs:
           41723
     (Ac) cDNA Polynucleotide Sequence
20
          - Pat. Appln. SEQ ID NO 39
          - Ceres seq id 1011954
        (B) Polypeptide Sequence
2.5
            - Pat. Appln. SEQ ID NO 40
            - Ceres seq id 1011955
            - Location of start within SEQ ID NO 39: at 1 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 40: at 40 aa.
30
         (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 41
            - Ceres seq id 1011956
            - Location of start within SEQ ID NO 39: at 107 nt.
35
          (Ba) Polypeptide Activities: Similar to DNAJ Protein
     Activities
      Maximum Length Sequence:
40
      related to:
     Clone IDs:
           41730
     Public Genomic DNA:
           qi No: 4678371
45
           Predicted Exons:
              SINGLE 48996 ... 49184 GENBANK
              SINGLE 48996 ... 49184 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 42
50
          - Ceres seq id 1011960
```

301

- Alternative transcription start site(s) located in SEQ ID NO 42:

20,22,25,43

- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 43
  - Ceres seq id 1011961
  - Location of start within SEQ ID NO 42: at 69 nt.
- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 44
  - Ceres seq id 1011962
  - Location of start within SEQ ID NO 42: at 78 nt.
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 45
  - Ceres seq id 1011963
  - Location of start within SEQ ID NO 42: at 90 nt.
- 20 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant specific gene, similar to serine rich protein

activities.

25

45

Maximum Length Sequence:

related to:

Clone IDs:

- 30 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 46
  - Ceres seq id 1014075
  - Alternative transcription start site(s) located in SEQ ID NO 46:
- 35 -3, -2, -1, 2, 3, 4, 5, 6, 7, 8, 10, 15, 103
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 47
    - Ceres seq\_id 1014076
- Location of start within SEQ ID NO 46: at 1 nt. 40
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 48
    - Ceres seq id 1014077
  - Location of start within SEQ ID NO 46: at 84 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 48: at 22 aa.
- (Ba) Polypeptide Activities: Arabidopsis specific gene, 50 dicot

302

specific gene, plant

```
specific gene.
```

Maximum Length Sequence:

5 related to:

Clone IDs:

37081

Public Genomic DNA:

gi No: 4159707

10 Predicted Exons:

INIT 15458 ... 15533 OCKHAMG-CDS TERM 15648 ... 15751 OCKHAMG-CDS

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 49
- Ceres seq id 1015865
  - Alternative transcription start site(s) located in SEQ ID NO 49:

-29

- 20 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 50
  - Ceres seq id 1015866
  - Location of start within SEQ ID NO 49: at 281 nt.
- 25 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 51
  - Ceres seq id 1015867
  - Location of start within SEQ ID NO 49: at 317 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 30 ID NO 51: at 22 aa.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 52
    - Ceres seq id 1015868
- 35 Location of start within SEQ ID NO 49: at 3 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 52: at 15  $\rm aa.$

(Ba) Polypeptide Activities: Arabidopsis specific

40 gene, dicot

specific gene, plant

specific gene.

Maximum Length Sequence:

45 related to:

Clone IDs:

29375

Public Genomic DNA:

gi No: 4159707

50 Predicted Exons:

INTR 15158 ... 15201 OCKHAMG-CDNA

303

INTR 15449 ... 15533 OCKHAMG-CDNA INTR 15648 ... 15947 OCKHAMG-CDNA

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 53
- Ceres seg id 1021371
- Alternative transcription start site(s) located in SEQ ID NO 53:

-74

- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 54
  - Ceres seq id 1021372
  - Location of start within SEQ ID NO 53: at 3 nt.
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 55
  - Ceres seq id 1021373
  - Location of start within SEQ ID NO 53: at 54 nt.
- 20 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 56
  - Ceres seq id 1021374
  - Location of start within SEQ ID NO 53: at 90 nt.
  - Location of Signal Peptide Cleavage Site within SEO
- 25 ID NO 56: at 22 aa.
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

30 specific gene.

Maximum Length Sequence: related to:

Clone IDs:

35 27064

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 57
  - Ceres seq id 1022578

40

5

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 58
  - Ceres seq id 1022579
  - Location of start within SEQ ID NO 57: at 1 nt.

45

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 59
  - Ceres seq id 1022580
  - Location of start within SEQ ID NO 57: at 157 nt.

50

(B) Polypeptide Sequence

304

```
- Pat. Appln. SEQ ID NO 60
```

- Ceres seq id 1022581
- Location of start within SEQ ID NO 57: at 10 nt.

5 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant

specific gene.

10 Maximum Length Sequence:

related to:

Clone IDs:

20

25

35

40

45

23773

Public Genomic DNA:

15 gi No: 6136357

Predicted Exons:

 INTR
 51972 ...
 51830 OCKHAMG-CDNA

 INTR
 51129 ...
 51055 OCKHAMG-CDNA

 INTR
 50948 ...
 50806 OCKHAMG-CDNA

 INTR
 50706 ...
 50608 OCKHAMG-CDNA

 INTR
 50326 ...
 50147 OCKHAMG-CDNA

gi No: 6143858

Predicted Exons:

INTR 51972 ... 51830 OCKHAMG-CDNA
INTR 51129 ... 51055 OCKHAMG-CDNA
INTR 50948 ... 50806 OCKHAMG-CDNA
INTR 50706 ... 50608 OCKHAMG-CDNA
INTR 50326 ... 50147 OCKHAMG-CDNA

- (Ac) cDNA Polynucleotide Sequence
- 30 Pat. Appln. SEQ ID NO 61
  - Ceres seq id 1024240
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 62
  - Ceres seq id 1024241
    - Location of start within SEQ ID NO 61: at 54 nt.
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 63
      - Ceres seq\_id 1024242
      - Location of start within SEQ ID NO 61: at 66 nt.
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 64
      - Ceres seq id 1024243
        - Location of start within SEQ ID NO 61: at 404 nt.

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

50 specific gene, plant specific gene.

```
Maximum Length Sequence:
      related to:
     Clone IDs:
 5
           2030
     Public Genomic DNA:
           gi No: 5441914
           Predicted Exons:
              SINGLE 4802 ... 4449
                                       OCKHAMG-CDS
           gi No: 5881519
10
           Predicted Exons:
              SINGLE 4802 ... 4449
                                        GENBANK
              SINGLE 4802 ... 4449
                                        OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
15
          - Pat. Appln. SEQ ID NO 65
          - Ceres seq id 1026562
          - Alternative transcription start site(s) located in SEQ
     ID NO 65:
20
            36, 41, 43, 44, 88, 90, 94, 102
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 66
            - Ceres seq id 1026563
25
            - Location of start within SEQ ID NO 65: at 1 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 66: at 51 aa.
        (B) Polypeptide Sequence
30
            - Pat. Appln. SEQ ID NO 67
            - Ceres seq id 1026564
            - Location of start within SEQ ID NO 65: at 109 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 67: at 15 aa.
35
          (Ba) Polypeptide Activities: Similar to cytochrome c
     oxidase II
                                         activities.
40
     Maximum Length Sequence:
      related to:
     Clone IDs:
           20097
     Public Genomic DNA:
45
           gi No: 5733893
           Predicted Exons:
              INTR
                     66948 ... 67429 OCKHAMG-CDNA
              SINGLE 66986 ... 67324 OCKHAMG-CDS
50
           qi No: 5748882
           Predicted Exons:
```

```
306
              INTR
                    41416 ...
                                40935
                                        OCKHAMG-CDNA
              SINGLE 41380 ... 41042 OCKHAMG-CDS
           gi No: 5801671
 5
           Predicted Exons:
                    51184 ... 50703
              INTR
                                       OCKHAMG-CDNA
              SINGLE 51148 ... 50810 OCKHAMG-CDS
          qi No: 5809708
10
           Predicted Exons:
                    41416 ...
              INTR
                                40935
                                       OCKHAMG-CDNA
              SINGLE 41380 ... 41042
                                        OCKHAMG-CDS
           gi No: 5836114
15
           Predicted Exons:
                    42487 ...
              INTR
                                42006
                                       OCKHAMG-CDNA
              SINGLE 42451 ... 42113 OCKHAMG-CDS
           gi No: 5923662
20
           Predicted Exons:
                     43106 ...
                                43587 OCKHAMG-CDNA
              INTR
              SINGLE 43144 ... 43482 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
25
          - Pat. Appln. SEQ ID NO 68
          - Ceres seq id 1026648
          - Alternative transcription start site(s) located in SEQ
     ID NO 68:
            -74, -72, -48, -42, 76
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 69
            - Ceres seq id 1026649
            - Location of start within SEQ ID NO 68: at 39 nt.
35
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 69: at 59 aa.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 70
40
            - Ceres seq id 1026650
            - Location of start within SEQ ID NO 68: at 78 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 70: at 46 aa.
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 71
            - Ceres seq id 1026651
            - Location of start within SEQ ID NO 68: at 81 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 71: at 45 aa.
50
```

307

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant

specific gene.

5

30

Maximum Length Sequence:

related to:

Clone IDs:

18274

- 10 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 72
  - Ceres seq id 1027881
  - Alternative transcription start site(s) located in SEQ ID NO 72:
- -8, -4, 2
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 73
    - Ceres seq id 1027882
- Location of start within SEQ ID NO 72: at 3 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 74
    - Ceres seq id 1027883
- 25 Location of start within SEQ ID NO 72: at 36 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 75
    - Ceres seg id 1027884
    - Location of start within SEQ ID NO 72: at 105 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 75: at 18 aa.
- (Ba) Polypeptide Activities: Arabidopsis specific
- 35 gene, dicot

specific gene, plant

specific gene.

Maximum Length Sequence:

40 related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 76
- 45 Ceres seq id 1381797
  - Alternative transcription start site(s) located in SEQ ID NO 76:
    - -41,18,19,22,26
- 50 (B) Polypeptide Sequence Pat. Appln. SEQ ID NO 77

308

- Ceres seq id 1381798
- Location of start within SEQ ID NO 76: at 70 nt.
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 78
  - Ceres seq id 1381799
  - Location of start within SEQ ID NO 76: at 82 nt.
- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 79
  - Ceres seq id 1381800
  - Location of start within SEQ ID NO 76: at 109 nt.

(Ba) Polypeptide Activities: Arabidopsis specific 15 gene, dicot specific gene, plant

specific gene.

Maximum Length Sequence:

20 related to:

5

10

Clone IDs:

31667

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 80
- 25 - Ceres seq id 1442747
  - Alternative transcription start site(s) located in SEQ ID NO 80:
    - 3, 5, 6, 11, 62, 64, 65, 66, 69, 70, 71, 72, 74
- 30 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 81
  - Ceres seq id 1442748
  - Location of start within SEQ ID NO 80: at 2 nt.
- 35 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 82
  - Ceres seq id 1442749
  - Location of start within SEQ ID NO 80: at 140 nt.
- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 83
  - Ceres seq id 1442750
  - Location of start within SEQ ID NO 80: at 224 nt.
- 45 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant

specific gene.

50 Maximum Length Sequence: related to:

309

Clone IDs:

270354

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 84
- 5 Ceres seq id 1459199
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 85
- Ceres seq id 1459200
  - Location of start within SEQ ID NO 84: at 2 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 86
- Ceres seq id 1459201
  - Location of start within SEQ ID NO 84: at 56 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 86: at 20 aa.
- 20 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 87
  - Ceres seq\_id 1459202
  - Location of start within SEQ ID NO 84: at 80 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 25 ID NO 87: at 17 aa.
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot
- 30 specific gene, plant specific gene.
- 35 Maximum Length Sequence:

related to:

Clone IDs:

12250

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 88
    - Ceres seq id 1565605
- Alternative transcription start site(s) located in SEQ ID NO 88:

-17

45

50

40

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 89
  - Ceres seq id 1565606
  - Location of start within SEQ ID NO 88: at 133 nt.

(B) Polypeptide Sequence

310

```
- Pat. Appln. SEQ ID NO 90
```

- Ceres seq id 1565607
- Location of start within SEQ ID NO 88: at 181 nt.
- 5 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

10 Maximum Length Sequence:

related to:

Clone IDs:

97883

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 91
  - Ceres seq id 1566686
  - Alternative transcription start site(s) located in SEQ ID NO 91:

58,61

20

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 92
  - Ceres seq id 1566687
  - Location of start within SEQ ID NO 91: at 137 nt.

25

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

30

Maximum Length Sequence:

related to:

Clone IDs:

- 35 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 93
  - Ceres seq id 1567367
  - Alternative transcription start site(s) located in SEQ ID NO 93:
- 40 -39,6,11,14,17,25,32,33,34,35,60
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 94
    - Ceres seq id 1567368
- 45 Location of start within SEQ ID NO 93: at 1 nt.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 95
    - Ceres seq id 1567369
- 50 Location of start within SEQ ID NO 93: at 2 nt.

311

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 96
            - Ceres seg id 1567370
            - Location of start within SEQ ID NO 93: at 65 nt.
 5
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
                                        specific gene, plant
     specific gene.
10
    Maximum Length Sequence:
     related to:
     Clone IDs:
           19528
15
     Public Genomic DNA:
           qi No: 6091770
           Predicted Exons:
                     21323 ...
              INIT
                                21390
                                        OCKHAMG-CDS
                     21467 ... 21564 OCKHAMG-CDS
              INTR
20
                     21995 ... 22027
              INTR
                                        OCKHAMG-CDS
                     22386 ... 22468
              TERM
                                        OCKHAMG-CDS
           gi No: 6102636
           Predicted Exons:
                   21323 ... 21390 OCKHAMG-CDS
              INIT
25
              INTR
                     21467 ... 21564 OCKHAMG-CDS
                     21995 ... 22027 OCKHAMG-CDS
              INTR
                     22386 ... 22468
              TERM
                                        OCKHAMG-CDS
           gi No: 6137907
           Predicted Exons:
30
                     21252 ... 21319 OCKHAMG-CDS
              INTT
              INTR
                     21396 ... 21493
                                        OCKHAMG-CDS
                     21924 ... 21956
              INTR
                                        OCKHAMG-CDS
              TERM
                     22315 ...
                                22397
                                        OCKHAMG-CDS
           gi No: 6437519
35
           Predicted Exons:
                     14107 ...
              INIT
                                14174
                                        OCKHAMG-CDS
                     14251 ...
              INTR
                                14348
                                        OCKHAMG-CDS
                     14779 ... 14811
              INTR
                                        OCKHAMG-CDS
                     15170 ... 15252
              TERM
                                        OCKHAMG-CDS
40
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 97
          - Ceres seq id 1570101
          - Alternative transcription start site(s) located in SEQ
     ID NO 97:
            9, 29, 30, 31, 32, 39, 40, 43, 44, 61, 62, 64, 65, 198
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 98
            - Ceres seq id 1570102
```

- Location of start within SEQ ID NO 97: at 116 nt.

312

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

5

Maximum Length Sequence:

related to:

Clone IDs:

42399

- 10 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 99
  - Ceres seq id 1571051
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 100
  - Ceres seq id 1571052
  - Location of start within SEQ ID NO 99: at 1 nt.
- 20 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 101
  - Ceres seq id 1571053
  - Location of start within SEQ ID NO 99: at 16 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 25 ID NO 101: at 30 aa.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 102
    - Ceres seq id 1571054
    - Location of start within SEQ ID NO 99: at 139 nt.

30

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

35

Maximum Length Sequence:

related to:

Clone IDs:

- 40 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 103
  - Ceres seq id 1571100
  - Alternative transcription start site(s) located in SEQ ID NO 103:
- 45 5,6
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 104
    - Ceres seq id 1571101
- 50 Location of start within SEQ ID NO 103: at 1 nt.

313

```
- Location of Signal Peptide Cleavage Site within SEQ
     ID NO 104: at 19 aa.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 105
5
            - Ceres seq id 1571102
            - Location of start within SEQ ID NO 103: at 82 nt.
             (Ba) Polypeptide Activities: Arabidopsis specific
     gene, dicot
10
                                        specific gene, plant
     specific gene.
    Maximum Length Sequence:
      related to:
15
    Clone IDs:
           39977
           14890
           34623
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 106
20
          - Ceres seq id 1665272
          - Alternative transcription start site(s) located in SEQ
     ID NO 106:
25
     3,15,32,33,34,35,37,38,39,40,41,42,43,44,45,46,47,48,49,50
            51,60,61,63,66,69,70,71,72,74,75,76,78,79,84,85
          - Clone 39977 starts at 35 and ends at 636 in cDNA.
          - Clone 14890 starts at 43 and ends at in cDNA.
          - Clone 34623 starts at 1 and ends at 598 in cDNA.
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 107
            - Ceres seq_id 1665273
            - Location of start within SEQ ID NO 106: at 90 nt.
35
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
                                         specific gene, plant
     specific gene.
40
     Maximum Length Sequence:
      related to:
     Clone IDs:
           24562
     (Ac) cDNA Polynucleotide Sequence
45
          - Pat. Appln. SEQ ID NO 108
          - Ceres seq id 1713895
          - Alternative transcription start site(s) located in SEQ
     ID NO 108:
```

-7,2,3,5,7,8,13,17,29,30,32,34

314

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 109
  - Ceres seq id 1713896
  - Location of start within SEQ ID NO 108: at 3 nt.

5

35

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 110
  - Ceres seq id 1713897
  - Location of start within SEQ ID NO 108: at 57 nt.
- 10 Location of Signal Peptide Cleavage Site within SEQ ID NO 110: at 20 aa.
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 111
- Ceres seq id 1713898
  - Location of start within SEQ ID NO 108: at 81 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 111: at 17 aa.
- 20 (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant

specific gene.

25 Maximum Length Sequence:

related to:

Clone IDs:

- (Ac) cDNA Polynucleotide Sequence
- Pat. Appln. SEQ ID NO 112
  - Ceres seq id 1923752
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 113
    - Ceres seq id 1923753
      - Location of start within SEQ ID NO 112: at 2 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 113: at 19  $\rm aa.$
- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 114
  - Ceres seq id 1923754
  - Location of start within SEQ ID NO 112: at 292 nt.
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 115
  - Ceres seq id 1923755
  - Location of start within SEQ ID NO 112: at 384 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 50 ID NO 115: at 27 aa.

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

315

specific gene, plant

specific gene, plant

specific gene.

5

Maximum Length Sequence:

related to:

Clone IDs:

150069

- 10 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 116
  - Ceres seq id 1976816
  - Alternative transcription start site(s) located in SEQ ID NO 116:
- 15 20,30,32

25

30

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 117
  - Ceres seq\_id 1976817
- 20 Location of start within SEQ ID NO 116: at 2 nt.
  - Location of Signal Peptide Cleavage Site within SEQ ID NO 117: at 61 aa.
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 118
      - Ceres seq id 1976818
      - Location of start within SEQ ID NO 116: at 3 nt.
    - (B) Polypeptide Sequence
      - Pat. Appln. SEQ ID NO 119
        - Ceres seq id 1976819
        - Location of start within SEQ ID NO 116: at 52 nt.
- (Ba) Polypeptide Activities: Arabidopsis specific gene, 35 dicot

specific gene.

Maximum Length Sequence:

40 related to:

Clone IDs:

109997

Public Genomic DNA:

gi No: 4263813

45 Predicted Exons:

INIT 82737 ... 82740 OCKHAMG-CDS TERM 83475 ... 83623 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 120
- 50 Ceres seq id 2025128

```
- Alternative transcription start site(s) located in SEQ
     ID NO 120:
            -5,5,61,88
          - Clone 109997 starts at 5 and ends at 566 in cDNA.
 5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 121
            - Ceres seq id 2025129
            - Location of start within SEQ ID NO 120: at 1 nt.
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 122
            - Ceres seq id 2025130
            - Location of start within SEQ ID NO 120: at 3 nt.
15
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 122: at 21 aa.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 123
20
            - Ceres seq id 2025131
            - Location of start within SEQ ID NO 120: at 316 nt.
             (Ba) Polypeptide Activities: Arabidopsis specific
    gene, dicot
25
                                        specific gene, plant
    specific gene.
    Maximum Length Sequence:
      related to:
    Clone IDs:
30
           41792
     Public Genomic DNA:
           qi No: 5762549
           Predicted Exons:
                    41272 ... 41749 OCKHAMG-CDNA
35
              INTR
           gi No: 5776585
           Predicted Exons:
                     55175 ... 55652 OCKHAMG-CDNA
              INTR
           qi No: 5801669
           Predicted Exons:
40
              INTR 97802 ... 98279 OCKHAMG-CDNA
           qi No: 5809709
           Predicted Exons:
              INTR
                    97803 ... 98280 OCKHAMG-CDNA
45
           qi No: 5932531
           Predicted Exons:
                    90652 ... 91129 OCKHAMG-CDNA
              INTR
           qi No: 6013612
           Predicted Exons:
50
              INTR
                     9143 ... 9620 OCKHAMG-CDNA
           gi No: 6016671
```

317 Predicted Exons: INTR 9266 ... 9743 OCKHAMG-CDNA gi No: 6041828 Predicted Exons: 5 INTR 9266 ... 9743 OCKHAMG-CDNA qi No: 6175159 Predicted Exons: INTR 92644 ... 92167 OCKHAMG-CDNA (Ac) cDNA Polynucleotide Sequence 10 - Pat. Appln. SEQ ID NO 124 - Ceres seg id 2025402 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 125 15 - Ceres seq id 2025403 - Location of start within SEQ ID NO 124: at 186 nt. (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 126 20 - Ceres seq id 2025404 - Location of start within SEQ ID NO 124: at 2 nt. (Ba) Polypeptide Activities: Similar to ribose 5phosphate isomerase 25 Maximum Length Sequence: related to: Clone IDs: 28572 Public Genomic DNA: 30 gi No: 4725940 Predicted Exons: 99197 ... 99167 OCKHAMG-CDS INIT 99060 ... 98960 INTR OCKHAMG-CDS 98863 ... 98753 OCKHAMG-CDS 35 INTR 98654 ... 97823 OCKHAMG-CDS INTR 97730 ... 97477 OCKHAMG-CDS INTR 96772 ... 96683 OCKHAMG-CDS INTR 96157 ... 96086 OCKHAMG-CDS TERM 40 qi No: 5823567 Predicted Exons: 99197 ... 99167 OCKHAMG-CDS INIT INTR 99060 ... 98960 OCKHAMG-CDS 98863 ... 98753 OCKHAMG-CDS INTR 45 INTR 98654 ... 97823 OCKHAMG-CDS 97730 ... 97477 OCKHAMG-CDS INTR 96772 ... INTR 96683 OCKHAMG-CDS 96157 ... 96086 OCKHAMG-CDS TERM (Ac) cDNA Polynucleotide Sequence 50 - Pat. Appln. SEQ ID NO 127

- Ceres seq id 2025479

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gi No: 6449509 Predicted Exons:

318 - Alternative transcription start site(s) located in SEQ ID NO 127: 1419 - Clone 28572 starts at 1419 and ends at 1955 in cDNA. (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 128 - Ceres seq id 2025480 - Location of start within SEQ ID NO 127: at 1 nt. (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 129 - Ceres seq id 2025481 - Location of start within SEQ ID NO 127: at 226 nt. (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 130 - Ceres seq id 2025482 - Location of start within SEQ ID NO 127: at 271 nt. (Ba) Polypeptide Activities: Similar Cell Wall Plasma Membrane disconnecting CLCT Protein activities, and proline rich protein activities. Maximum Length Sequence: Public Genomic DNA: gi No: 6:34247 Predicted Exons: INTR 46663 ... 46055 OCKHAMG-CDNA SINGLE 46579 ... 46103 OCKHAMG-CDS (Ac) cDNA Polynucleotide Sequence - Pat. Appln. SEQ ID NO 131 - Ceres seq id 2032963 (B) Polypeptide Sequence - Pat. Appln. SEQ ID NO 132 - Ceres seq id 2032964 - Location of start within SEQ ID NO 131: at 85 nt. (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot specific gene, plant specific gene. Maximum Length Sequence: Public Genomic DNA:

```
319
             INIT 32110 ... 32118 OCKHAMG-CDS
             INTR 32363 ... 32538 OCKHAMG-CDS
             INTR 33006 ... 33084 OCKHAMG-CDS
             TERM 33182 ... 33247 OCKHAMG-CDS
 5
          gi No: 6453848
          Predicted Exons:
             INIT 34795 ... 34803 OCKHAMG-CDS
             INTR
                   35048 ... 35223 OCKHAMG-CDS
                    35691 ... 35769 OCKHAMG-CDS
             INTR
10
             TERM 35867 ... 35932 OCKHAMG-CDS
          gi No: 6456153
          Predicted Exons:
             INIT
                   32305 ... 32313 OCKHAMG-CDS
             INTR
                    32558 ... 32733 OCKHAMG-CDS
             INTR 33201 ... 33279 OCKHAMG-CDS
15
             TERM 33377 ... 33442 OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
         - Pat. Appln. SEQ ID NO 133
20
         - Ceres seq id 2033706
       (B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 134
           - Ceres seq id 2033707
25
           - Location of start within SEQ ID NO 133: at 1 nt.
       (B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 135
           - Ceres seq id 2033708
30
           - Location of start within SEQ ID NO 133: at 34 nt.
       (B) Polypeptide Sequence
           - Pat. Appln. SEQ ID NO 136
           - Ceres seq id 2033709
35
           - Location of start within SEQ ID NO 133: at 160 nt.
         (Ba) Polypeptide Activities: Arabidopsis specific gene,
    dicot
                                      specific gene, plant
40
    specific gene.
    Maximum Length Sequence:
    Public Genomic DNA:
          gi No: 4512646
45
          Predicted Exons:
             INTR 43130 ... 43387 OCKHAMG-CDNA
             INTR 43479 ... 43638 OCKHAMG-CDNA
             INIT 43224 ... 43387 OCKHAMG-CDS
50
             TERM 43479 ... 43494 OCKHAMG-CDS
```

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320
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 137
          - Ceres seq id 2043118
 5
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 138
            - Ceres seq id 2043119
            - Location of start within SEQ ID NO 137: at 65 nt.
            - Location of Signal Peptide Cleavage Site within SEQ
     ID NO 138: at 22 aa.
10
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 139
            - Ceres seq id 2043120
15
            - Location of start within SEQ ID NO 137: at 1 nt.
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 140
            - Ceres seq id 2043121
20
            - Location of start within SEQ ID NO 137: at 95 nt.
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
     dicot
                                        specific gene, plant
25
     specific gene.
     Maximum Length Sequence:
     Public Genomic DNA:
           qi No: 3985934
30
           Predicted Exons:
                    40094 ... 40516 OCKHAMG-CDNA
              INTR
                    40861 ... 41275 OCKHAMG-CDNA
              INTR
              INIT 40233 ... 40516 OCKHAMG-CDS
35
                    40861 ... 41077 OCKHAMG-CDS
              TERM
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 141
          - Ceres seq id 2047214
40
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 142
            - Ceres seq id 2047215
            - Location of start within SEQ ID NO 141: at 140 nt.
45
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 143
            - Ceres seq id 2047216
            - Location of start within SEQ ID NO 141: at 224 nt.
```

(B) Polypeptide Sequence

321

- Pat. Appln. SEQ ID NO 144
- Ceres seq id 2047217
- Location of start within SEQ ID NO 141: at 153 nt.
- Location of Signal Peptide Cleavage Site within SEQ
- 5 ID NO 144: at 18 aa.
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

10 specific gene.

Maximum Length Sequence: Public Genomic DNA:

gi No: 3985958

15 Predicted Exons:

INTR	8598	 8696	OCKHAMG-CDNA
INTR	8960	 9035	OCKHAMG-CDNA
INTR	9202	 9513	OCKHAMG-CDNA
INTR	9628	 9669	OCKHAMG-CDNA

20

30

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- INIT 8997 ... 9035 OCKHAMG-CDS INTR 9202 ... 9513 OCKHAMG-CDS TERM 9628 ... 9822 OCKHAMG-CDS
- 25 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 145
  - Ceres seq id 2047438
  - (B) Polypeptide Sequence
    - Pat. Appln. SEQ ID NO 146
    - Ceres seq id 2047439
    - Location of start within SEQ ID NO 145: at 137 nt.

Maximum Length Sequence:

35 Public Genomic DNA:

gi No: 4757403 Predicted Exons:

INTR 35281 ... 34781 OCKHAMG-CDNA

40 SINGLE 35235 ... 35005 OCKHAMG-CDS

- (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 147
  - Ceres seq id 2049056

(B) Polymen

- (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 148
  - Ceres seq id 2049057
  - Location of start within SEQ ID NO 147: at 47 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 148: at 22 aa.

322

```
(B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 149
           - Ceres seg id 2049058
 5
           - Location of start within SEQ ID NO 147: at 59 nt.
            - Location of Signal Peptide Cleavage Site within SEO
     ID NO 149: at 18 aa.
    Maximum Length Sequence:
10
     Public Genomic DNA:
          gi No: 4538972
           Predicted Exons:
                    32316 ...
              INTR
                               32561 OCKHAMG-CDNA
             INTR 32643 ... 32715
                                       OCKHAMG-CDNA
                    33489 ... 33697 OCKHAMG-CDNA
15
             INTR
             INTR
                    33789 ... 33935 OCKHAMG-CDNA
             INTR 34034 ... 34365 OCKHAMG-CDNA
             INIT 32328 ... 32561 OCKHAMG-CDS
             INTR 32643 ... 32715 OCKHAMG-CDS
20
             INTR 33489 ... 33697 OCKHAMG-CDS
                    33789 ... 33935
             INTR
                                       OCKHAMG-CDS
             TERM
                    34034 ... 34159 OCKHAMG-CDS
25
     (Ac) cDNA Polynucleotide Sequence
         - Pat. Appln. SEQ ID NO 150
         - Ceres seq id 2050386
        (B) Polypeptide Sequence
30
           - Pat. Appln. SEQ ID NO 151
           - Ceres seq id 2050387
           - Location of start within SEQ ID NO 150: at 1 nt.
        (B) Polypeptide Sequence
35
           - Pat. Appln. SEQ ID NO 152
           - Ceres seq id 2050388
           - Location of start within SEQ ID NO 150: at 13 nt.
        (B) Polypeptide Sequence
40
           - Pat. Appln. SEQ ID NO 153
           - Ceres seq id 2050389
           - Location of start within SEQ ID NO 150: at 151 nt.
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
45
    dicot
                                       specific gene, plant
     specific gene.
    Maximum Length Sequence:
50
    Public Genomic DNA:
```

gi No: 4878038

```
323
          Predicted Exons:
                    22640 ...
                               22509 OCKHAMG-CDNA
             INTR
                    22412 ...
             INTR
                               22340 OCKHAMG-CDNA
                    22122 ...
                               21728 OCKHAMG-CDNA
             INTR
5
                    22674 ...
             INTR
                               22509
                                       OCKHAMG-CDNA
                    22412 ...
                              22340
             INTR
                                       OCKHAMG-CDNA
                    22122 ...
                               21766
                                       OCKHAMG-CDNA
             INTR
10
                    22587 ...
                               22509 OCKHAMG-CDS
             TINI
             INTR
                    22412 ... 22340
                                       OCKHAMG-CDS
                    22122 ...
                               21852
             TERM
                                       OCKHAMG-CDS
          qi No: 6143825
          Predicted Exons:
15
             INTR
                    22639 ...
                               22508
                                       OCKHAMG-CDNA
             INTR
                    22411 ...
                               22339
                                       OCKHAMG-CDNA
                    22121 ...
                               21727
             INTR
                                       OCKHAMG-CDNA
                    22673 ...
                               22508
             INTR
                                       OCKHAMG-CDNA
20
             INTR
                    22411 ...
                               22339
                                       OCKHAMG-CDNA
                    22121 ...
             INTR
                               21765
                                       OCKHAMG-CDNA
                    22586 ...
                               22508
                                       OCKHAMG-CDS
             INIT
                    22411 ...
                               22339
                                       OCKHAMG-CDS
             INTR
25
             TERM
                    22121 ...
                               21851
                                       OCKHAMG-CDS
     (Ac) cDNA Polynucleotide Sequence
          - Pat. Appln. SEQ ID NO 154
         - Ceres seq id 2053353
30
        (B) Polypeptide Sequence
            - Pat. Appln. SEQ ID NO 155
            - Ceres seq id 2053354
           - Location of start within SEQ ID NO 154: at 88 nt.
35
          (Ba) Polypeptide Activities: Arabidopsis specific gene,
    dicot
                                       specific gene, plant
     specific gene.
40
    Maximum Length Sequence:
     Public Genomic DNA:
          gi No: 4115352
45
           Predicted Exons:
                    29289 ...
                               29287
              INIT
                                       OCKHAMG-CDS
              INTR
                    27681 ...
                               27485
                                       OCKHAMG-CDS
                    27312 ...
                               27193
              INTR
                                       OCKHAMG-CDS
                    26979 ... 26916
              INTR
                                       OCKHAMG-CDS
50
                    26654 ...
                                     OCKHAMG-CDS
             TERM
                               26646
```

324

(Ac) cDNA Polynucleotide Sequence

- Pat. Appln. SEQ ID NO 156
- Ceres seq id 2055693
- 5 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 157
  - Ceres seq id 2055694
  - Location of start within SEQ ID NO 156: at 1 nt.
- 10 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 158
  - Ceres seq id 2055695
  - Location of start within SEQ ID NO 156: at 46 nt.
- 15 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 159
  - Ceres seq id 2055696
  - Location of start within SEQ ID NO 156: at 2 nt.
  - Location of Signal Peptide Cleavage Site within SEQ
- 20 ID NO 159: at 28 aa.
  - (Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

25 specific gene.

Maximum Length Sequence:

Public Genomic DNA:

gi No: 4406790

30 Predicted Exons:

INTR 81640 ... 81247 OCKHAMG-CDNA

SINGLE 81603 ... 81367 OCKHAMG-CDS

- 35 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 160
  - Ceres seq id 2056405
- 40 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 161
  - Ceres seq id 2056406
  - Location of start within SEQ ID NO 160: at 38 nt.
- 45 (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 162
  - Ceres seq id 2056407
  - Location of start within SEQ ID NO 160: at 56 nt.
- 50 (B) Polypeptide Sequence
   Pat. Appln. SEQ ID NO 163

325

- Ceres seq id 2056408
- Location of start within SEQ ID NO 160: at 177 nt.
- Location of Signal Peptide Cleavage Site within SEQ ID NO 163: at 14 aa.

5

(Ba) Polypeptide Activities: Arabidopsis specific gene, dicot

specific gene, plant

specific gene.

10

25

Maximum Length Sequence:
 related to:

Clone IDs:

233233

- 15 (Ac) cDNA Polynucleotide Sequence
  - Pat. Appln. SEQ ID NO 164
  - Ceres seq id 2065747
  - (B) Polypeptide Sequence
- 20 Pat. Appln. SEQ ID NO 165
  - Ceres seq id 2065748
  - Location of start within SEQ ID NO 164: at 114 nt.
  - (B) Polypeptide Sequence
  - Pat. Appln. SEQ ID NO 166
    - Ceres seq id 2065749
    - Location of start within SEQ ID NO 164: at 279 nt.
- (Ba) Polypeptide Activities: Arabidopsis specific gene, 30 dicot specific gene, plant

specific gene.

```
SEQ TABLE 1
      (2) INFORMATION FOR SEQ ID NO:1:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 601 base pairs
 5
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
10
                (A) NAME/KEY: -
                (B) LOCATION: 1..601
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007546
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
      qacqaacaqq ccacqcqtqa ctacatacat aatttqqttt ctctttttt ttttqttctt
                                                                             60
15
      cttgttcctc tttggttttt gacattggtt ctggagaaga tgagttcggt agaaccagac
                                                                             120
      atggaagatc tqttccaqqa qaaqaaqcqt qtcaqqaatc ctctcqttcc tctcqqtqca
                                                                             180
      cttatgaccg cgggagtgct cacggctggg ttgattagtt tcagaagagg caattctcag
                                                                             240
      ttgggtcagg ttttgatgag agctagggtg gtcgtccagg gtgctactgt cgctttaatg
      gttggaaccg gttattacta cggtgataat ccgtggaaga agtgagctcc aactacttgt
                                                                             360
20
      tcgttttcac ataattqaqa tqacactcgg qaaatctgcg ggcaaaggtg tgtgtgctct
                                                                             420
      cqctctqaqt qtaqaaatat qcqqtcttqc taqqatttaq taaqqqtaca ttacatqatt
                                                                             480
      tgaataattt gaagaacgaa caagtatcgt tgtatcttga tcaattgatt agattgattt
                                                                             540
      aaggaaattc tccqaaaacc atctttactc tatctgtcaa aatctcatta tctttctgcc
                                                                             600
25
      (2) INFORMATION FOR SEQ ID NO:2:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 114 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..114
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007547
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
      Asp Glu Gln Ala Thr Arg Asp Tyr Ile His Asn Leu Val Ser Leu Phe
                                           10
      Phe Phe Val Leu Leu Val Pro Leu Trp Phe Leu Thr Leu Val Leu Glu
40
                                       25
      Lys Met Ser Ser Val Glu Pro Asp Met Glu Asp Leu Phe Gln Glu Lys
                                  40
      Lys Arg Val Arg Asn Pro Leu Val Pro Leu Gly Ala Leu Met Thr Ala
                              55
45
      Gly Val Leu Thr Ala Gly Leu Ile Ser Phe Arg Arg Gly Asn Ser Gln
                          70
                                               75
      Leu Gly Gln Val Leu Met Arg Ala Arg Val Val Val Gln Gly Ala Thr
                      85
                                           90
      Val Ala Leu Met Val Gly Thr Gly Tyr Tyr Gly Asp Asn Pro Trp
50
                                       105
                  100
      Lys Lys
      (2) INFORMATION FOR SEQ ID NO:3:
           (i) SEQUENCE CHARACTERISTICS:
55
                (A) LENGTH: 81 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
60
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..81
```

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(D) OTHER INFORMATION: / Ceres Seq. ID 1007548
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
     Met Ser Ser Val Glu Pro Asp Met Glu Asp Leu Phe Gln Glu Lys Lys
                                         10
 5
     Arg Val Arg Asn Pro Leu Val Pro Leu Gly Ala Leu Met Thr Ala Gly
                                     2.5
     Val Leu Thr Ala Gly Leu Ile Ser Phe Arg Arg Gly Asn Ser Gln Leu
                                 40
     Gly Gln Val Leu Met Arg Ala Arg Val Val Gln Gly Ala Thr Val
10
                            55
                                                 60
     Ala Leu Met Val Gly Thr Gly Tyr Tyr Tyr Gly Asp Asn Pro Trp Lys
                         70
                                             75
     Lys
15
     (2) INFORMATION FOR SEQ ID NO:4:
          (i) SEOUENCE CHARACTERISTICS:
               (A) LENGTH: 74 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
20
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..74
25
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007549
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
      Met Glu Asp Leu Phe Gln Glu Lys Lys Arg Val Arg Asn Pro Leu Val
                                         10
      Pro Leu Gly Ala Leu Met Thr Ala Gly Val Leu Thr Ala Gly Leu Ile
30
                 2.0
                                     25
      Ser Phe Arg Arg Gly Asn Ser Gln Leu Gly Gln Val Leu Met Arg Ala
                                 40
      Arg Val Val Gln Gly Ala Thr Vai Ala Leu Met Val Gly Thr Gly
35
      Tyr Tyr Tyr Gly Asp Asn Pro Trp Lys Lys
                         70
      (2) INFORMATION FOR SEO ID NO:5:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 970 base pairs
40
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
45
                (A) NAME/KEY: -
                (B) LOCATION: 1..970
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007583
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
      aacgccgctc atttttaggg tttcagatat aggcagattc tcacatttgc aaacatgacg
                                                                           60
50
      120
                                                                          180
      gaggttctta caagagagct agcagaggat ggttactctg gtgttgaggt tagggttact
                                                                          240
      cctatgagga ctgagattat catcagagct acacgtactc agaatgttct cggtgagaag
                                                                          300
      gggaggagaa ttagggaatt gacttccctt gttcagaaga gattcaagtt tccagttgac
      agtgttgage tttatgccga gaaggttaac aacagaggte tetgtgccat cgctcagget
                                                                          360
55
                                                                          420
      qagtctttac qttacaaqct tcttqqtqqt ctcqctqttc qtaqqqcttq ctatqqtqtq
                                                                          480
      ttgaggtttg ttatggagag tggagctaag ggatgcgagg ttatcgtgag tggaaagctt
      cgtgctgcca gagccaagtc tatgaagttc aaagatggct acatggtgtc atctggtcaa
                                                                          540
      ccaactaaqq aatacataqa ctctqcaqtq aqacatqttt tqcttaqaca aggtqtgttq
                                                                          600
      ggaatcaagg tgaaggttat gcttgattgg gaccctaagg gcatatcagg accaaagaca
                                                                          660
60
      ccattgcctg atgttgtgat cattcattct cctaaagaag aagaggccat ctacgcacct
                                                                          720
      gctcaggttg ctgccccggc tgctctcgta gcagatgcac cactcacagc cgtagattac
                                                                          780
      cctgcgatga tcccagtcgc ctaaaagaat cctttgtact agtcgatgtg attttgtttt
                                                                          840
```

328

caactgttct tgctcttttg acagattata gcttctcgta tcttcgaatc agacacgttt 900 cctctgttct tattcgaaaa ttatctcaag tttccattgt tgaaaaagcc ctttttagtt 960 tcataaatcg (2) INFORMATION FOR SEQ ID NO:6: 5 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 249 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 10 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..249 (D) OTHER INFORMATION: / Ceres Seq. ID 1007584 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: Met Thr Thr Gln Ile Ser Lys Lys Arg Lys Phe Val Ala Asp Gly Val 5 10 Phe Tyr Ala Glu Leu Asn Glu Val Leu Thr Arg Glu Leu Ala Glu Asp 20 25 20 Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile 40 Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg 55 Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys Arg Phe Lys Phe Pro 25 70 75 Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val Asn Asn Arg Gly Leu 90 Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly 105 30 Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu 120 125 Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala 135 140 Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Val Ser Ser 35 150 155 Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ser Ala Val Arg His Val Leu 170 165 Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys Val Met Leu Asp Trp 185 40 Asp Pro Lys Gly Ile Ser Gly Pro Lys Thr Pro Leu Pro Asp Val Val 195 200 Ile Ile His Ser Pro Lys Glu Glu Glu Ala Ile Tyr Ala Pro Ala Gln 215 220 Val Ala Ala Pro Ala Ala Leu Val Ala Asp Ala Pro Leu Thr Ala Val 45 230 235 Asp Tyr Pro Ala Met Ile Pro Val Ala 245 (2) INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS: 50 (A) LENGTH: 206 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 55 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..206 (D) OTHER INFORMATION: / Ceres Seq. ID 1007585 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 60 Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu 10 Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys

```
25
                  20
                                                           30
     Arg Phe Lys Phe Pro Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val
                                  40
     Asn Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr
 5
     Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu
                                              75
     Arg Phe Val Met Glu Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser
                                          90
10
      Gly Lys Leu Arg Ala Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly
                  100
                                      105
                                                           110
      Tyr Met Val Ser Ser Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ser Ala
              115
                                  120
      Val Arg His Val Leu Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys
15
                              135
                                                   140
      Val Met Leu Asp Trp Asp Pro Lys Gly Ile Ser Gly Pro Lys Thr Pro
                          150
                                               155
      Leu Pro Asp Val Val Ile Ile His Ser Pro Lys Glu Glu Glu Ala Ile
                      165
                                           170
20
      Tyr Ala Pro Ala Gln Val Ala Ala Pro Ala Ala Leu Val Ala Asp Ala
                                      185
      Pro Leu Thr Ala Val Asp Tyr Pro Ala Met Ile Pro Val Ala
                                  200
              195
      (2) INFORMATION FOR SEQ ID NO:8:
25
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 768 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..768
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008148
35
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
      aaaaaaaccc tagttttttg ctctctcgaa gttccgtgcg actctctgct cgagcaagat
                                                                              60
      ctcctaccga tacaatgatt atctcagagg ctaaccgcaa agaaatctgc aagtacctct
                                                                             120
      tcaaagaagg agtttgcttt gcgaagaagg atttcaatct tgctaagcat ccgttgatcg
                                                                             180
      atgtcccgaa cctacaagtg attaagctta tgcagagttt caaatccaag gagtatgtta
                                                                             240
40
      gagagacatt tgcctggatg cattactatt ggtttttgac caatgaaggg atcgagttct
                                                                             300
      tgagaactta tcttaatctt ccatctgatg ttgttcctgc tactttgaag aagtcagcta
                                                                             360
      agectggtgg tegtecattt ggtggeceae etggtgateg etcaagagga eetegeeatg
                                                                             420
      aaggaggaga ccqtcccagg tttggtgacc gtgatgggta ccqtqcaggt cctcgagctg
                                                                             480
      gtggtgagtt tggaggtgaa aagggtggag ctcctgcaga ttaccagcca tctttccaag
                                                                             540
45
      qaaqtqqqcq tqqttttqqc cqtqqtqctq qtqqctacaq cqcaqctqca ccatctqqtt
                                                                             600
      caggtttqcc ttqaaaaaqa aatqtcttta qqtqacaqta aqaccatqqa qqaqttttca
                                                                             660
      gctttaaatt ttgcttttgt aattcagatt ccggaatcct tcataatctc tatctgagtt
                                                                             720
      tagttttgtt gttgaatcaa acatccgatt taaagttatg ttcattcc
      (2) INFORMATION FOR SEQ ID NO:9:
50
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 203 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
55
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..203
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1008149
60
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
      Lys Asn Pro Ser Phe Leu Leu Ser Arg Ser Ser Val Arg Leu Ser Ala
                       5
                                           10
```

330 Arg Ala Arg Ser Pro Thr Asp Thr Met Ile Ile Ser Glu Ala Asn Arg .25 Lys Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly Val Cys Phe Ala Lys 40 Lys Asp Phe Asn Leu Ala Lys His Pro Leu Ile Asp Val Pro Asn Leu 55 Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly 10 90 Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro Ser Asp Val Val Pro 105 Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly 120 15 Pro Pro Gly Asp Arg Ser Arg Gly Pro Arg His Glu Gly Gly Asp Arg 135 140 Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Ala Gly Pro Arg Ala Gly 150 155 Gly Glu Phe Gly Gly Glu Lys Gly Gly Ala Pro Ala Asp Tyr Gln Pro 20 170 165 Ser Phe Gln Gly Ser Gly Arg Gly Phe Gly Arg Gly Ala Gly Gly Tyr 185 180 Ser Ala Ala Pro Ser Gly Ser Gly Leu Pro 25 (2) INFORMATION FOR SEQ ID NO:10: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 179 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..179 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1008150 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: Met Ile Ile Ser Glu Ala Asn Arg Lys Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Ala Lys His 40 25 Pro Leu Ile Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser 40 Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr 45 Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu 75 70 Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys 90 Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Ser Arg Gly 50 105 Pro Arg His Glu Gly Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly 120 Tyr Arg Ala Gly Pro Arg Ala Gly Gly Glu Phe Gly Gly Glu Lys Gly 55 Gly Ala Pro Ala Asp Tyr Gln Pro Ser Phe Gln Gly Ser Gly Arg Gly 150 155 Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser 170 165

(2) INFORMATION FOR SEQ ID NO:11:
 (i) SEQUENCE CHARACTERISTICS:

Gly Leu Pro

```
(A) LENGTH: 134 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..134
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008151
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
      Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp
                                          10
      Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg
                                      25
15
      Thr Tyr Leu Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys
                                  40
      Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg
      Ser Arg Gly Pro Arg His Glu Gly Gly Asp Arg Pro Arg Phe Gly Asp
20
                          70
                                              75
      Arg Asp Gly Tyr Arg Ala Gly Pro Arg Ala Gly Gly Glu Phe Gly Gly
                                           90
                      85
      Glu Lys Gly Gly Ala Pro Ala Asp Tyr Gln Pro Ser Phe Gln Gly Ser
                 . 100
                                      105
25
      Gly Arg Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Pro
              115
                                  120
      Ser Gly Ser Gly Leu Pro
          130
      (2) INFORMATION FOR SEQ ID NO:12:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 995 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..995
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008334
40
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:
      atatetaaaa gaeteaaaaa eetategtea tttatettea acaggeggaa gaacggegat
                                                                              60
      ggccgccatt acagetetca etetecgete teetgttttt ettetteete categteegt
                                                                             120
                                                                             180
      cacaagccct agattccatg gcttcaccaa tcaaccacca ccagctcgtc tcttctttcc
      tettaaecce tteeettete tateaateea aaaecetaaa teeateegaa ttteegeate
                                                                              240
45
      cgcttcgccg ataacaacac caatcctcca gacggagaag tcaacagctc ggtcatcgac
                                                                              300
                                                                              360
      actcaccggg tccactcggt ctctcgctac tctcgcggct ttagcaatcg ctgtaaccag
      agtectaget cagaaactet etetegeaat ceagaettea agtecegtaa tegeegaegg
                                                                              420
      gttacgattc tctctcagta ccgccggacc tgtcttcttc gcgtctctcc gggatcgtcc
                                                                              480
      tccgggatac ttgaacacgc cgctgacggt tgttgcggtg gggataaaga agtggctaga
                                                                              540
50
      catttacagt ggggtattga tggttagggt tttgctgagt tggttcccta atatcccttg
                                                                              600
      ggaaagacag cetttgtetg ceattagaga tetetgtgat cettaettga atetetteag
                                                                              660
      aaacatcatt cctcctatct tcgatacgct tgatgttagt ccattgcttg ctttcgcggt
                                                                              720
      tcttggtaca cttggatcga ttgttcatgg cagcactggg tagaaattgg aagacttgag
                                                                             780
      ctaatgqttt tqttqaaacq ttcaattqqq aaatttctqc aacttqtttc ccaaqattaq
                                                                             840
55
      qqaaaaatca qaatttqatq ctqtqttqqa tqctcqatqa qaqttqtqaq ctttatcqtt
                                                                             900
      tttgttgttc tctttttgac tttgattctg aataatgaga ttttgggttg ttgtttatct
                                                                             960
      cctaattaca tgttcagact acagtttatt gatgg
      (2) INFORMATION FOR SEQ ID NO:13:
            (i) SEQUENCE CHARACTERISTICS:
60
                 (A) LENGTH: 253 amino acids
                 (B) TYPE: amino acid
```

(C) STRANDEDNESS:

```
(D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
5
               (B) LOCATION: 1..253
               (D) OTHER INFORMATION: / Ceres Seq. ID 1008335
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
     Tyr Leu Lys Asp Ser Lys Thr Tyr Arg His Leu Ser Ser Thr Gly Gly
                                        10
10
     Arg Thr Ala Met Ala Ala Ile Thr Ala Leu Thr Leu Arg Ser Pro Val
                                     25
     Phe Leu Leu Pro Pro Ser Ser Val Thr Ser Pro Arg Phe His Gly Phe
                                 40
     Thr Asn Gln Pro Pro Pro Ala Arg Leu Phe Phe Pro Leu Asn Pro Phe
15
                             55
     Pro Ser Leu Ser Ile Gln Asn Pro Lys Ser Ile Arg Ile Ser Ala Ser
                         70
                                             75
     Ala Ser Pro Ile Thr Thr Pro Ile Leu Gln Thr Glu Lys Ser Thr Ala
                                         90
20
     Arg Ser Ser Thr Leu Thr Gly Ser Thr Arg Ser Leu Ala Thr Leu Ala
                 100
                                     105
     Ala Leu Ala Ile Ala Val Thr Arg Val Leu Ala Gln Lys Leu Ser Leu
                                 120
     Ala Ile Gln Thr Ser Ser Pro Val Ile Ala Asp Gly Leu Arg Phe Ser
25
                             135
     Leu Ser Thr Ala Gly Pro Val Phe Phe Ala Ser Leu Arg Asp Arg Pro
                         150
                                             155
      Pro Gly Tyr Leu Asn Thr Pro Leu Thr Val Val Ala Val Gly Ile Lys
                                        170
                     165
30
     Lys Trp Leu Asp Ile Tyr Ser Gly Val Leu Met Val Arg Val Leu Leu
                                     185
      Ser Trp Phe Pro Asn Ile Pro Trp Glu Arg Gln Pro Leu Ser Ala Ile
                                200
      Arg Asp Leu Cys Asp Pro Tyr Leu Asn Leu Phe Arg Asn Ile Ile Pro
                  215
                                                220
35
      Pro Ile Phe Asp Thr Leu Asp Val Ser Pro Leu Leu Ala Phe Ala Val
                        230
                                            235
      Leu Gly Thr Leu Gly Ser Ile Val His Gly Ser Thr Gly
                     245
40
      (2) INFORMATION FOR SEQ ID NO:14:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 234 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..234
50
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008336
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
      Met Ala Ala Ile Thr Ala Leu Thr Leu Arg Ser Pro Val Phe Leu Leu
                                         10
      Pro Pro Ser Ser Val Thr Ser Pro Arg Phe His Gly Phe Thr Asn Gln
55
                                     25
      Pro Pro Pro Ala Arg Leu Phe Pro Leu Asn Pro Phe Pro Ser Leu
                                 40
      Ser Ile Gln Asn Pro Lys Ser Ile Arg Ile Ser Ala Ser Ala Ser Pro
                              55
60
      Ile Thr Thr Pro Ile Leu Gln Thr Glu Lys Ser Thr Ala Arg Ser Ser
                                             75
                          70
      Thr Leu Thr Gly Ser Thr Arg Ser Leu Ala Thr Leu Ala Ala Leu Ala
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333
                     8.5
                                          90
      Ile Ala Val Thr Arg Val Leu Ala Gln Lys Leu Ser Leu Ala Ile Gln
                                     105 110
      Thr Ser Ser Pro Val Ile Ala Asp Gly Leu Arg Phe Ser Leu Ser Thr
5
                                 120
                                                      125
      Ala Gly Pro Val Phe Phe Ala Ser Leu Arg Asp Arg Pro Pro Gly Tyr
                             135
                                                  140
      Leu Asn Thr Pro Leu Thr Val Val Ala Val Gly Ile Lys Lys Trp Leu
                          150
                                              155
10
      Asp Ile Tyr Ser Gly Val Leu Met Val Arg Val Leu Leu Ser Trp Phe
                      165
                                          170
      Pro Asn Ile Pro Trp Glu Arg Gln Pro Leu Ser Ala Ile Arg Asp Leu
                                      185
      Cys Asp Pro Tyr Leu Asn Leu Phe Arg Asn Ile Ile Pro Pro Ile Phe
15
                                  200
                                                      205
      Asp Thr Leu Asp Val Ser Pro Leu Leu Ala Phe Ala Val Leu Gly Thr
                              215
                                                  220
      Leu Gly Ser Ile Val His Gly Ser Thr Gly
                          230
20
      (2) INFORMATION FOR SEQ ID NO:15:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 520 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..520
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008701
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:
      acacataacc ctcttggaaa gagtctcaac acttgcagag aaaaagaaga aggaagatcc
                                                                             60
      cggaaaatgg caacggcgat tgtacgttca gctctttccc gagcagcgac tcgcgcagct
                                                                             120
      ccgaagacat ccgtcgctcc taagcgaaac ttttcctctt ccgccggcca tgacgatgct
                                                                             180
35
      tatgaagctg cgaagtggga gaagataact tatctgggta ttgctagttg cactgctcta
                                                                             240
                                                                             300
      gctgtctatg ttttatccaa gggccatcat cccggcgaag accctcctgc ctatccgcat
                                                                             360
      atgcacatcc gcaacaagga gtttccttgg ggtccggatg gtctgtttga ggtgaagcac
      aacaaagagc actgagtctt gcgtggtcat aataacgtct tcttggttta tttgaaaggc
                                                                             420
      taaaatgttt taccgtattt gttctcaccg tttgtcaacg attttctact ccaatctctt
                                                                             480
40
      ttcttttgtt gggaaataaa agttaatact ttgcttggtc
      (2) INFORMATION FOR SEQ ID NO:16:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 124 amino acids
                (B) TYPE: amino acid
45
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
50
                (B) LOCATION: 1..124
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008702
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:
      Thr His Asn Pro Leu Gly Lys Ser Leu Asn Thr Cys Arg Glu Lys Glu
                                          10
55
      Glu Gly Arg Ser Arg Lys Met Ala Thr Ala Ile Val Arg Ser Ala Leu
                                       25
      Ser Arg Ala Ala Thr Arg Ala Ala Pro Lys Thr Ser Val Ala Pro Lys
                                  40
      Arg Asn Phe Ser Ser Ser Ala Gly His Asp Asp Ala Tyr Glu Ala Ala
60
                              55
                                                   60
      Lys Trp Glu Lys Ile Thr Tyr Leu Gly Ile Ala Ser Cys Thr Ala Leu
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334 Ala Val Tyr Val Leu Ser Lys Gly His His Pro Gly Glu Asp Pro Pro 85 · 90 Ala Tyr Pro His Met His Ile Arg Asn Lys Glu Phe Pro Trp Gly Pro 100 105 110 5 Asp Gly Leu Phe Glu Val Lys His Asn Lys Glu His 115 120 (2) INFORMATION FOR SEQ ID NO:17: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 102 amino acids 10 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 15 (A) NAME/KEY: peptide (B) LOCATION: 1..102 (D) OTHER INFORMATION: / Ceres Seq. ID 1008703 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17: Met Ala Thr Ala Ile Val Arg Ser Ala Leu Ser Arg Ala Ala Thr Arg 20 5 10 Ala Pro Lys Thr Ser Val Ala Pro Lys Arg Asn Phe Ser Ser Ser 25 Ala Gly His Asp Asp Ala Tyr Glu Ala Ala Lys Trp Glu Lys Ile Thr 40 25 Tyr Leu Gly Ile Ala Ser Cys Thr Ala Leu Ala Val Tyr Val Leu Ser 55 Lys Gly His His Pro Gly Glu Asp Pro Pro Ala Tyr Pro His Met His 70 75 Ile Arg Asn Lys Glu Phe Pro Trp Gly Pro Asp Gly Leu Phe Glu Val 30 90 8.5 Lys His Asn Lys Glu His 100 (2) INFORMATION FOR SEQ ID NO:18: (i) SEQUENCE CHARACTERISTICS: 35 (A) LENGTH: 68 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 40 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..68 (D) OTHER INFORMATION: / Ceres Seq. ID 1008704 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18: 45 His Ile Thr Leu Leu Glu Arg Val Ser Thr Leu Ala Glu Lys Lys 10 Lys Glu Asp Pro Gly Lys Trp Gln Arg Arg Leu Tyr Val Gln Leu Phe 25 Pro Glu Gln Arg Leu Ala Gln Leu Arg Arg His Pro Ser Leu Leu Ser 50 40 Glu Thr Phe Pro Leu Pro Pro Ala Met Thr Met Leu Met Lys Leu Arg 55 Ser Gly Arg Arg 55 (2) INFORMATION FOR SEQ ID NO:19: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 662 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 60 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE:

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335
                (A) NAME/KEY: -
                (B) LOCATION: 1..662
                (D) OTHER INFORMATION: / Ceres Seq. ID 1009003
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:
 5
      cattttttct ttctctctct cctccctaag caaaactaaa acaagctatg gctggtatgc
                                                                             60
      ttcccggagt tgagtgtgca aggaggcggc gcttccacgg tggtgctcct ccgattgaat
                                                                             120
      cctcgaacac agcttctgtg gcggctgcgg cgggacacgt ctggacacgg cgaccatcgt
                                                                             180
      tototottta cactaccaat catgagagoc accaggocca tgtotootto toggagagaa
                                                                             240
      gtgttaggaa taaatcttat ggagaagaca acgatgagaa acttgacgga gcagccaaag
                                                                             300
10
      aggcaaagca gaggcttaac aagcggctga gaatcccacc acgtacaagt tcaggcaaaa
                                                                             360
      tggtaaagac aaagggaata aattggagca aggaaagggt aaacctctcg gggacttacc
                                                                             420
      gaccgaggtg gtcgggttaa agaagagccg aggaaggttg atggaatggt tcaagcggcg
                                                                             480
      agttagggaa caacaagatt gtgctatatg tctagaccgg ttcaagaagg gtgagacctt
                                                                             540
      ggtacaccta ccatgtgccc ataagtttca ctccatatgc ttattgcctt ggctagacac
                                                                             600
15
      taatgtttat tgcccatatt gtagaactga tatttggaat taaatgttat atttttgatg
                                                                             660
      (2) INFORMATION FOR SEO ID NO:20:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 130 amino acids
20
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
25
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..130
                (D) OTHER INFORMATION: / Ceres Seq. ID 1009004
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:
      Met Ala Gly Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Phe
30
                                          10
      His Gly Gly Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala
                  20
                                      25
      Ala Ala Ala Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr
                                  40
35
      Thr Thr Asn His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arq
      Ser Val Arg Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp
                          70
                                               75
      Gly Ala Ala Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile
40
      Pro Pro Arg Thr Ser Ser Gly Lys Met Val Lys Thr Lys Gly Ile Asn
                                      105
                  100
                                                           110
      Trp Ser Lys Glu Arg Val Asn Leu Ser Gly Thr Tyr Arg Pro Arg Trp
              115
                                  120
45
      Ser Gly
          130
      (2) INFORMATION FOR SEQ ID NO:21:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 127 amino acids
50
                 (B) TYPE: amino acid
                (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
55
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..127
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1009005
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:
      Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Phe His Gly Gly
60
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10

Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala Ala Ala Ala

336 Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr Thr Thr Asn 40 4.5 His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arg Ser Val Arg 55 60 5 Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp Gly Ala Ala 70 75 Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile Pro Pro Arg Thr Ser Ser Gly Lys Met Val Lys Thr Lys Gly Ile Asn Trp Ser Lys 10 100 105 Glu Arg Val Asn Leu Ser Gly Thr Tyr Arg Pro Arg Trp Ser Gly 120 (2) INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 504 base pairs 15 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 20 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..504 (D) OTHER INFORMATION: / Ceres Seq. ID 1009345 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22: 25 aagaaacaaa ggtgtcaaga agaaggatca acatattaat tgactaaaaa tgaatgcatg gatgataatc ttgttggtga tttgtgtcgc tgtggtggtg gagcaatcag aggctcgcaa 120 aggtcgaaag tatttaaatc caggcgtgct tgaccggtgt cgtggtccta atcctccagc 180 gggatgtcat cctcacaatt cccaccacaa acctcgcgtc cctgttcaca attatagtcg 240 tggttgtagt agaattaccc ggtgcagacg agatgcctag gtttaacctc acatgacgac 300 30 gacgtccctt tatattgatc cttcttctac atgagacttg caatgatatt atatattcat 360 atatatacac attacgctgt atatgataat tccagttgat taatatataa atcttactga 420 tttcaattca atacacaaga tcctgtaact gaaataattt tgtccatgtt ttgtgcttat 480 aataatggta tgactcttat tatt (2) INFORMATION FOR SEQ ID NO:23: 35 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 76 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 40 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..76 (D) OTHER INFORMATION: / Ceres Seq. ID 1009346 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23: Met Asn Ala Trp Met Ile Ile Leu Leu Val Ile Cys Val Ala Val Val 5 10 Val Glu Gln Ser Glu Ala Arg Lys Gly Arg Lys Tyr Leu Asn Pro Gly 25 50 Val Leu Asp Arg Cys Arg Gly Pro Asn Pro Pro Ala Gly Cys His Pro 40 His Asn Ser His His Lys Pro Arg Val Pro Val His Asn Tyr Ser Arg Gly Cys Ser Arg Ile Thr Arg Cys Arg Arg Asp Ala 55 70 (2) INFORMATION FOR SEQ ID NO:24: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 72 amino acids (B) TYPE: amino acid 60 (C) STRANDEDNESS: (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

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(ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..72
                (D) OTHER INFORMATION: / Ceres Seq. ID 1009347
 5
          (xi) SEQUENCE DESCRIPTION: SEO ID NO:24:
     Met Ile Ile Leu Leu Val Ile Cys Val Ala Val Val Glu Gln Ser
                      5
                                          10
     Glu Ala Arg Lys Gly Arg Lys Tyr Leu Asn Pro Gly Val Leu Asp Arg
                  20
                                      2.5
10
      Cys Arg Gly Pro Asn Pro Pro Ala Gly Cys His Pro His Asn Ser His
              35
                                  40
      His Lys Pro Arg Val Pro Val His Asn Tyr Ser Arg Gly Cys Ser Arg
                              55
      Ile Thr Arg Cys Arg Arg Asp Ala
15
                          70
      (2) INFORMATION FOR SEQ ID NO:25:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 967 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..967
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010140
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:
      attitattic tictccaacc attictcaatc cictctctcc aatggcticc tiatccttcg
                                                                              60
      tttcctcttc tcaccttacg ctacgcactc cttctattgc cctacgcagc actggctctt
                                                                             120
30
      ctcctcgaac ctccgtttca ttctccgtca aggctcaatc cgtcgcgctt tcacaggacg
                                                                             180
      atttgaagaa gctcgcggcg gagaaagctg tggaggcaat taaacctggg atggttctag
                                                                             240
      ggctcggaac cggatccacc gcagctttcg ctgttgatca gatcgggaaa ctactctctt
                                                                             300
                                                                             360
      ccggtgaact ctacgatatt gtcggtatcc caacgtcgaa acgaacggag gaacaagcac
                                                                             420
      ggtcgttagg gattcctctt gttgggttag atacacatcc gagaatcgat ctcgctattg
35
      acggagcaga cgaggtagat ccgaatcttg atttagtcaa aggtcgtgga ggtgctcttc
                                                                             480
      tccgtgagaa aatggtggaa gctgtggctg acaagtttat tgttgtggct gatgatacca
                                                                             540
                                                                             600
      aactcgttac aggactcggt ggaagtggat tagctatgcc ggtggaagtt gttcaattct
                                                                             660
      gctggaactt taatttgatt agattgcaag agctcttcaa ggaatttgga tgtgaatcaa
                                                                             720
      agcttagagt tgatggtgat ggcaagcctt atgtgactga taacagtaat tacattattg
40
      atttgtattt taagactcct ttgaaggatg gattcgctgc ggctaaagag attgggaagt
                                                                             780
      ttcaaqqagt qqtqqaqcat qqtctqtttc tcqqaatqqc tacttctgtc attatcgctq
                                                                             840
                                                                             900
      qaaaqaatgg cqttqaagtt atgaccaagt qaggttaaaa gttcaatctt tctcttcttc
      tttgattgac aaatttttct cctttgtaaa attatcaatt gagtttttat gtgagatcat
                                                                             960
      ttcaatc
45
      (2) INFORMATION FOR SEQ ID NO:26:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 289 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
50
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..289
55
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1010141
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:
      Phe Ile Ser Ser Pro Thr Ile Leu Asn Pro Leu Ser Pro Met Ala Ser
                                           10
      Leu Ser Phe Val Ser Ser Ser His Leu Thr Leu Arg Thr Pro Ser Ile
60
                  20
                                      25
                                                           30
      Ala Leu Arg Ser Thr Gly Ser Ser Pro Arg Thr Ser Val Ser Phe Ser
               35
                                   40
                                                       45
```

```
Val Lys Ala Gln Ser Val Ala Leu Ser Gln Asp Asp Leu Lys Lys Leu
                         55
     Ala Ala Glu Lys Ala Val Glu Ala Ile Lys Pro Gly Met Val Leu Gly
                        70
 5
     Leu Gly Thr Gly Ser Thr Ala Ala Phe Ala Val Asp Gln Ile Gly Lys
                   85
                                       90
     Leu Leu Ser Ser Gly Glu Leu Tyr Asp Ile Val Gly Ile Pro Thr Ser
                100 105
     Lys Arg Thr Glu Glu Gln Ala Arg Ser Leu Gly Ile Pro Leu Val Gly
10
                                120
                                                   125
     Leu Asp Thr His Pro Arg Ile Asp Leu Ala Ile Asp Gly Ala Asp Glu
                           135
                                                140
     Val Asp Pro Asn Leu Asp Leu Val Lys Gly Arg Gly Gly Ala Leu Leu
                        150
                                          155
15
     Arg Glu Lys Met Val Glu Ala Val Ala Asp Lys Phe Ile Val Val Ala
                                        170
     Asp Asp Thr Lys Leu Val Thr Gly Leu Gly Gly Ser Gly Leu Ala Met
                                    185
     Pro Val Glu Val Val Gln Phe Cys Trp Asn Phe Asn Leu Ile Arg Leu
20
                                200
     Gln Glu Leu Phe Lys Glu Phe Gly Cys Glu Ser Lys Leu Arg Val Asp
                            215
                                                220
     Gly Asp Gly Lys Pro Tyr Val Thr Asp Asn Ser Asn Tyr Ile Ile Asp
                         230
                                            235
25
     Leu Tyr Phe Lys Thr Pro Leu Lys Asp Gly Phe Ala Ala Ala Lys Glu
                                        250
      Ile Gly Lys Phe Gln Gly Val Val Glu His Gly Leu Phe Leu Gly Met
                                    265
     Ala Thr Ser Val Ile Île Ala Gly Lys Asn Gly Val Glu Val Met Thr
30
                                 280
      Lys
      (2) INFORMATION FOR SEQ ID NO:27:
          (i) SEQUENCE CHARACTERISTICS:
35
               (A) LENGTH: 276 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..276
               (D) OTHER INFORMATION: / Ceres Seq. ID 1010142
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:
45
      Met Ala Ser Leu Ser Phe Val Ser Ser Ser His Leu Thr Leu Arg Thr
                                        10
      Pro Ser Ile Ala Leu Arg Ser Thr Gly Ser Ser Pro Arg Thr Ser Val
                                   25
                 20
      Ser Phe Ser Val Lys Ala Gln Ser Val Ala Leu Ser Gln Asp Asp Leu
50
                                40
      Lys Lys Leu Ala Ala Glu Lys Ala Val Glu Ala Ile Lys Pro Gly Met
                            55
      Val Leu Gly Leu Gly Thr Gly Ser Thr Ala Ala Phe Ala Val Asp Gln
                         70
                                            75
55
      Ile Gly Lys Leu Ser Ser Gly Glu Leu Tyr Asp Ile Val Gly Ile
                                         90
      Pro Thr Ser Lys Arg Thr Glu Glu Gln Ala Arg Ser Leu Gly Ile Pro
                                    105
      Leu Val Gly Leu Asp Thr His Pro Arg Ile Asp Leu Ala Ile Asp Gly
60
                                120
      Ala Asp Glu Val Asp Pro Asn Leu Asp Leu Val Lys Gly Arg Gly Gly
```

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339
    Ala Leu Leu Arg Glu Lys Met Val Glu Ala Val Ala Asp Lys Phe Ile
            150 155 160
    Val Val Ala Asp Asp Thr Lys Leu Val Thr Gly Leu Gly Gly Ser Gly
                      170
                 165
5
    Leu Ala Met Pro Val Glu Val Val Gln Phe Cys Trp Asn Phe Asn Leu
                   185
     Ile Arg Leu Gln Glu Leu Phe Lys Glu Phe Gly Cys Glu Ser Lys Leu
           195 200
                                205
     Arg Val Asp Gly Asp Gly Lys Pro Tyr Val Thr Asp Asn Ser Asn Tyr
10
     210 215
                                         220
     Ile Ile Asp Leu Tyr Phe Lys Thr Pro Leu Lys Asp Gly Phe Ala Ala
                     230
                                      235
     Ala Lys Glu Ile Gly Lys Phe Gln Gly Val Val Glu His Gly Leu Phe
                  245 250
15
     Leu Gly Met Ala Thr Ser Val Ile Ile Ala Gly Lys Asn Gly Val Glu
                       265
     Val Met Thr Lys
           275
     (2) INFORMATION FOR SEO ID NO:28:
20
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 213 amino acids
             (B) TYPE: amino acid
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
25
        (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..213
             (D) OTHER INFORMATION: / Ceres Seq. ID 1010143
30
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:
     Met Val Leu Gly Leu Gly Thr Gly Ser Thr Ala Ala Phe Ala Val Asp
                                   10
     Gln Ile Gly Lys Leu Leu Ser Ser Gly Glu Leu Tyr Asp Ile Val Gly
               20
                                25
35
     Ile Pro Thr Ser Lys Arg Thr Glu Glu Gln Ala Arg Ser Leu Gly Ile
                          40
     Pro Leu Val Gly Leu Asp Thr His Pro Arg Ile Asp Leu Ala Ile Asp
                        55
     Gly Ala Asp Glu Val Asp Pro Asn Leu Asp Leu Val Lys Gly Arg Gly
40
                                      75
     Gly Ala Leu Leu Arg Glu Lys Met Val Glu Ala Val Ala Asp Lys Phe
                                    90
     Ile Val Val Ala Asp Asp Thr Lys Leu Val Thr Gly Leu Gly Gly Ser
                                105
                                       110
45
     Gly Leu Ala Met Pro Val Glu Val Val Gln Phe Cys Trp Asn Phe Asn
                            120 125
     Leu Ile Arg Leu Gln Glu Leu Phe Lys Glu Phe Gly Cys Glu Ser Lys
                         135
                                          140
     Leu Arg Val Asp Gly Asp Gly Lys Pro Tyr Val Thr Asp Asn Ser Asn
50
                     150
                                      155
     Tyr Ile Ile Asp Leu Tyr Phe Lys Thr Pro Leu Lys Asp Gly Phe Ala
                  165
                                   170
     Ala Ala Lys Glu Ile Gly Lys Phe Gln Gly Val Val Glu His Gly Leu
              180 185
55
     Phe Leu Gly Met Ala Thr Ser Val Ile Ile Ala Gly Lys Asn Gly Val
           195
                       200
     Glu Val Met Thr Lys
        210
     (2) INFORMATION FOR SEQ ID NO:29:
60
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 623 base pairs
              (B) TYPE: nucleic acid
```

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(C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
 5
                (A) NAME/KEY: -
                (B) LOCATION: 1..623
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010217
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:
      qaaaaccctt aagctaagca aatttcaaag caattcaaaa accctagcct ctctctttt
10
      atctctcaca acaaaatctt tqaaatggcg atctctaagg cttccattgt tgttctcatg
                                                                             120
      atggtgatta totocgtogt tgcatoggca cagtotgagg caccagcacc aagtoctact
                                                                             180
      totggatota gtgcgatoto agogtottto gtgtcagotg tgcagttato accggcggag
                                                                             240
      agggaagcaa gggttttgag gtatagagag aagaggaaga atcggaaatt tgagaagacg
                                                                             300
      attaggtatg cgtcgcgtaa agcttacgct gagatgaggc cgagaatcaa aggacgtttt
                                                                             360
15
      gctaagagaa cagattcgag agagaatgat ggtggagacg tcggagttta ttgcggattc
                                                                             420
      ggtgttgttc cgagtttctg atatttcccg gttaaagaaa catggtagta gtatatcggt
                                                                             480
      taatagcaga ttaaggataa ttaaaactaa aatgtttcct gatttaatca ggggctaagt
                                                                             540
      tattgttaga agattgttgt tttttgaatt aattctcaca attgtatcct aaattatata
                                                                             600
      gtttacttaa tgttgaacta atg
20
      (2) INFORMATION FOR SEQ ID NO:30:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 118 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..118
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010218
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:
      Met Ala Ile Ser Lys Ala Ser Ile Val Val Leu Met Met Val Ile Ile
                      5
                                           10
      Ser Val Val Ala Ser Ala Gln Ser Glu Ala Pro Ala Pro Ser Pro Thr
35
                                       25
                                                           30
      Ser Gly Ser Ser Ala Ile Ser Ala Ser Phe Val Ser Ala Val Gln Leu
                                  4 0
                                                       45
      Ser Pro Ala Glu Arg Glu Ala Arg Val Leu Arg Tyr Arg Glu Lys Arg
                                                   60
                              55
40
      Lys Asn Arg Lys Phe Glu Lys Thr Ile Arg Tyr Ala Ser Arg Lys Ala
                                               75
                          70
      Tyr Ala Glu Met Arg Pro Arg Ile Lys Gly Arg Phe Ala Lys Arg Thr
                                           90
      Asp Ser Arg Glu Asn Asp Gly Gly Asp Val Gly Val Tyr Cys Gly Phe
45
                  100
                                       105
      Gly Val Val Pro Ser Phe
              115
      (2) INFORMATION FOR SEQ ID NO:31:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 107 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..107
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1010219
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:
60
      Met Met Val Ile Ile Ser Val Val Ala Ser Ala Gln Ser Glu Ala Pro
                                           10
```

Ala Pro Ser Pro Thr Ser Gly Ser Ser Ala Ile Ser Ala Ser Phe Val

341

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25
                                                          30
                  20
      Ser Ala Val Gln Leu Ser Pro Ala Glu Arg Glu Ala Arg Val Leu Arg
                                 4.0
      Tyr Arg Glu Lys Arg Lys Asn Arg Lys Phe Glu Lys Thr Ile Arg Tyr
 5
                              55
                                                  60
      Ala Ser Arg Lys Ala Tyr Ala Glu Met Arg Pro Arg Ile Lys Gly Arg
                          70
                                              75
      Phe Ala Lys Arg Thr Asp Ser Arg Glu Asn Asp Gly Gly Asp Val Gly
10
      Val Tyr Cys Gly Phe Gly Val Val Pro Ser Phe
                  100
      (2) INFORMATION FOR SEQ ID NO:32:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 106 amino acids
15
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
20
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..106
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010220
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:
      Met Val Ile Ile Ser Val Val Ala Ser Ala Gln Ser Glu Ala Pro Ala
25
                      5
                                           10
      Pro Ser Pro Thr Ser Gly Ser Ser Ala Ile Ser Ala Ser Phe Val Ser
      Ala Val Gln Leu Ser Pro Ala Glu Arg Glu Ala Arg Val Leu Arg Tyr
30
      Arg Glu Lys Arg Lys Asn Arg Lys Phe Glu Lys Thr Ile Arg Tyr Ala
                              55
      Ser Arg Lys Ala Tyr Ala Glu Met Arg Pro Arg Ile Lys Gly Arg Phe
                          70
                                              75
      Ala Lys Arg Thr Asp Ser Arg Glu Asn Asp Gly Gly Asp Val Gly Val
35
                      85
                                           90
      Tyr Cys Gly Phe Gly Val Val Pro Ser Phe
                  100
      (2) INFORMATION FOR SEQ ID NO:33:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 568 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
45
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..568
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010302
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:
50
      aaaacaaatc attcaaaqac atacaaaata attgagtttt ttttttaatt agaaacaaat
                                                                              60
      ggggttgagt ggtgttcttc atgtggaggt tgaggttaag tctccggctg aaaagttctg
                                                                             120
      ggtagccctc ggcgacggca tcaatctctt ccccaaagct ttccctaacg actacaaaac
                                                                             180
      catccaagtt ctagccggcg acggcaacgc tcctggctcc attcgcctca ttacttatgg
                                                                             240
      agaaggatct ccactggtga agatatcggc ggagaggatc gaagcagtgg atttggagaa
55
      caaaagcatg tcatacagca tcattggcgg cgaaatgttg gagtactaca aaacgttcaa
      aggaaccatc acceptattc ctaaqaacqq tqqcaqcctt ctqaaatggt ctggtgagtt
                                                                             420
      tgagaagacc gcccatgaga ttgatgaccc acacgtcatc aaggactttg ctgtcaagaa
                                                                             480
      cttcaaagag atagatgagt atctccttaa gcaaactagt gcctaacact agaaccttta
                                                                            540
      aattatataa gagggttcga tcgtctct
60
      (2) INFORMATION FOR SEQ ID NO:34:
           (i) SEQUENCE CHARACTERISTICS:
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(A) LENGTH: 155 amino acids

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(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..155
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010303
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:
10
     Met Gly Leu Ser Gly Val Leu His Val Glu Val Glu Val Lys Ser Pro
                      5
                                          10
     Ala Glu Lys Phe Trp Val Ala Leu Gly Asp Gly Ile Asn Leu Phe Pro
                                      25
      Lys Ala Phe Pro Asn Asp Tyr Lys Thr Ile Gln Val Leu Ala Gly Asp
15
                                  40
      Gly Asn Ala Pro Gly Ser Ile Arg Leu Ile Thr Tyr Gly Glu Gly Ser
                              55
      Pro Leu Val Lys Ile Ser Ala Glu Arg Ile Glu Ala Val Asp Leu Glu
                          70
                                               75
20
      Asn Lys Ser Met Ser Tyr Ser Ile Ile Gly Gly Glu Met Leu Glu Tyr
                                           90
      Tyr Lys Thr Phe Lys Gly Thr Ile Thr Val Ile Pro Lys Asn Gly Gly
                  100
                                      105
      Ser Leu Leu Lys Trp Ser Gly Glu Phe Glu Lys Thr Ala His Glu Ile
25
                                  120
      Asp Asp Pro His Val Ile Lys Asp Phe Ala Val Lys Asn Phe Lys Glu
                              135
                                                   140
      Ile Asp Glu Tyr Leu Leu Lys Gln Thr Ser Ala
                          150
30
      (2) INFORMATION FOR SEQ ID NO:35:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 531 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
35
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..531
40
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010815
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:
      gcgccgcccg aacccgaaga aqaactctct cttgtaaacc tgcgatctga atcttcccgg
                                                                              60
      cggtgaaaaa tggttgcgga gaagaaggcg aagaagtccc atgaaggaat caacagcaga
                                                                             120
                                                                             180
      ttagetettg tgatgaagag tggaaagtac actettggat acaagtetgt teteaaatee
45
                                                                              240
      cttcgcagct ccaaaggtaa gctgatattg atatctagca attgcccacc gttgagaagg
      tcagagattg aatactatgc gatgcttgct aaagttggag ttcatcgcta caatggcaac
                                                                              300
      aatgttgatt tgggtactgc ttgtggtaaa tacttccgag tttcttgcct cagcatcgtt
                                                                              360
                                                                             420
      gatcctggtg attccgacat catcaagaca cttcctggag atcagtgatt ctgattttga
      tgattttgcc atgttgttct ctatgcttga ttatgctttg tgttcttcaa agacttaatg
                                                                              480
50
      ttagatttct ttggtttccc gacttcctct taatggttat aattaattac t
      (2) INFORMATION FOR SEQ ID NO:36:
            (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 112 amino acids
                (B) TYPE: amino acid
55
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
60
                (B) LOCATION: 1..112
                (D) OTHER INFORMATION: / Ceres Seq. ID 1010816
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:
```

343 Met Val Ala Glu Lys Lys Ala Lys Lys Ser His Glu Gly Ile Asn Ser 10 Arg Leu Ala Leu Val Met Lys Ser Gly Lys Tyr Thr Leu Gly Tyr Lys 25 5 Ser Val Leu Lys Ser Leu Arg Ser Ser Lys Gly Lys Leu Ile Leu Ile Ser Ser Asn Cys Pro Pro Leu Arg Arg Ser Glu Ile Glu Tyr Tyr Ala Met Leu Ala Lys Val Gly Val His Arg Tyr Asn Gly Asn Asn Val Asp 10 Leu Gly Thr Ala Cys Gly Lys Tyr Phe Arg Val Ser Cys Leu Ser Ile 90 85 Val Asp Pro Gly Asp Ser Asp Ile Ile Lys Thr Leu Pro Gly Asp Gln 100 105 15 (2) INFORMATION FOR SEQ ID NO:37: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 91 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..91 25 (D) OTHER INFORMATION: / Ceres Seq. ID 1010817 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37: Met Lys Ser Gly Lys Tyr Thr Leu Gly Tyr Lys Ser Val Leu Lys Ser 5 10 Leu Arg Ser Ser Lys Gly Lys Leu Ile Leu Ile Ser Ser Asn Cys Pro 30 25 Pro Leu Arg Arg Ser Glu Ile Glu Tyr Tyr Ala Met Leu Ala Lys Val 40 Gly Val His Arg Tyr Asn Gly Asn Asn Val Asp Leu Gly Thr Ala Cys 55 35 Gly Lys Tyr Phe Arg Val Ser Cys Leu Ser Ile Val Asp Pro Gly Asp 70 Ser Asp Ile Ile Lys Thr Leu Pro Gly Asp Gln 85 (2) INFORMATION FOR SEQ ID NO:38: 40 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 73 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 45 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..73 (D) OTHER INFORMATION: / Ceres Seq. ID 1010818 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38: Met Arg Cys Leu Leu Lys Leu Glu Phe Ile Ala Thr Met Ala Thr Met 10 Leu Ile Trp Val Leu Leu Val Val Asn Thr Ser Glu Phe Leu Ala Ser 20 25 55 Ala Ser Leu Ile Leu Val Ile Pro Thr Ser Ser Arg His Phe Leu Glu 40 45 Ile Ser Asp Ser Asp Phe Asp Asp Phe Ala Met Leu Phe Ser Met Leu 55 Asp Tyr Ala Leu Cys Ser Ser Lys Thr 60 70 (2) INFORMATION FOR SEQ ID NO:39: (i) SEQUENCE CHARACTERISTICS:

```
(A) LENGTH: 494 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..494
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011437
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:
     acaatcggaa gaaaggagaa gacgaagatg tcgttggtat ggctggaagc gatgttgcct
                                                                           60
     ctcggaatca tcggtgggat gctctgtatc atgggcaatt ctcagtacta catccacaaa
     gcttatcatg gccgtcctaa gcacatcggc cacgatgaat gggatgtttc tatggaaaga
     cgcgacaaga aagtcgtcga gaaagctgca gctccttcct catgattcgc tttatctctt
15
     ttgtgttcct caggggctta aggtgaccac ttgtggtgac aaataaagtg cattccagaa
     gaagaagaag ctgggggatc tagtactttc attcccattt gattttcctt ggacatatta
     aagctttcag aaatcagacc tcaataacat ttggtttatc aatatttctc tattcgtgat
     attttgtatg cttttaagtt gcatgataca ctgaactact ctcttgctgt gtgagtgaat
     aaatgaatat ctgt
20
     (2) INFORMATION FOR SEO ID NO:40:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 94 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
25
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..94
30
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011438
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:
     Gln Ser Glu Glu Arg Arg Arg Arg Cys Arg Trp Tyr Gly Trp Lys
     1 5
                                        10
                                                            15
     Arg Cys Cys Leu Ser Glu Ser Ser Val Gly Cys Ser Val Ser Trp Ala
35
     Ile Leu Ser Thr Thr Ser Thr Lys Leu Ile Met Ala Val Leu Ser Thr
                                 40
      Ser Ala Thr Met Asn Gly Met Phe Leu Trp Lys Asp Ala Thr Arg Lys
                             55
40
     Ser Ser Arg Lys Leu Gln Leu Leu Pro His Asp Ser Leu Tyr Leu Phe
                                             75
      Cys Val Pro Gln Gly Leu Lys Val Thr Thr Cys Gly Asp Lys
                     85
      (2) INFORMATION FOR SEQ ID NO:41:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011439
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:
      Thr Ile Gly Arg Lys Glu Lys Thr Lys Met Ser Leu Val Trp Leu Glu
                                         10
     Ala Met Leu Pro Leu Gly Ile Ile Gly Gly Met Leu Cys Ile Met Gly
                                    25
60
      Asn Ser Gln Tyr Tyr Ile His Lys Ala Tyr His Gly Arg Pro Lys His
                                 40
      Ile Gly His Asp Glu Trp Asp Val Ser Met Glu Arg Arg Asp Lys Lys
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WO 00/40695 PCT/US00/00466 345 55 60 Val Val Glu Lys Ala Ala Ala Pro Ser Ser 70 (2) INFORMATION FOR SEQ ID NO:42: 5 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 65 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 10 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..65 (D) OTHER INFORMATION: / Ceres Seq. ID 1011440 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42: Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile Ile Gly 10 5 Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His Lys Ala 20 25 20 Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp Val Ser 40 45 Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Pro Ser 55 Ser 25 65 (2) INFORMATION FOR SEQ ID NO:43: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1011 base pairs (B) TYPE: nucleic acid 30 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -35 (B) LOCATION: 1..1011 (D) OTHER INFORMATION: / Ceres Seq. ID 1011616 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43: ctgtttcttt ttgcttcctt agacttctcc gtcgattaac ccttgccgag ggtcattttt ttttcttgaa ttttttcacc aagtgttcgt tcacggcatt gcttgtcctc cagtatgagt 120 40 atctccggtg ctgcagttgg ttcaggcaga aatttaagaa gagcggtaga gtttgggaaa 180 actcatgtgg ttaggcctaa agggaaacat caagcaacta ttgtctggtt acatgggctt 240 ggggacaatg gctcgagctg gtcccagctt ttggagaccc ttccccttcc aaatatcaaa 300 tggatttgcc cgactgctcc ttctcaacca ataagtttat ttggtggttt tccctccaca 360 gcttggtttg atgttgtgga catcaatgaa gatggacctg atgatatgga aggattggat 420 45 gtggctgctg cacatgttgc aaatctgttg tcgaatgagc ctgctgacat taaattaggt 480 gttggaggat tcagcatggg tgcggcgaca tctctatatt ctgcaacttg ttttgctctc 540 ggtaaatatg gaaatggcaa tccataccct atcaatttaa gcgcaatcat aggcttaagc 600 ggctggcttc cttgtgcaaa gacattggct ggcaaactag aagaggaaca gatcaagaac 660 cgagctgcat cgttacccat tgttgtctgt catggaaaag ctgatgatgt ggtaccgttc 720 50 aagtttgggg agaaatcttc acaggctttg ctttcaaatg ggtttaagaa ggtgaccttc 780 aaaccttaca gtgcacttgg tcaccacaca atcccacagg agttggatga gttgtgcgca 840 tggttgacat ccacgctcag cctcgaaggt tgatacttcc tatgatgtag ctttctgatg 900 aaaacccttc aactcttgag agtttgattg aattggatgt ttcaggattt cacaatgttt 960 tcattggaat atttgtgtaa gacacattcg tctagtagca aatcttctcc c 55 (2) INFORMATION FOR SEQ ID NO:44: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 252 amino acids (B) TYPE: amino acid (C) STRANDEDNESS:

(ii) MOLECULE TYPE: peptide
(ix) FEATURE:

(D) TOPOLOGY: linear

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(A) NAME/KEY: peptide
               (B) LOCATION: 1..252
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011617
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:
 5
     Met Ser Ile Ser Gly Ala Ala Val Gly Ser Gly Arg Asn Leu Arg Arg
                                       10
     Ala Val Glu Phe Gly Lys Thr His Val Val Arg Pro Lys Gly Lys His
                                   25
     Gln Ala Thr Ile Val Trp Leu His Gly Leu Gly Asp Asn Gly Ser Ser
10
                               40
     Trp Ser Gln Leu Leu Glu Thr Leu Pro Leu Pro Asn Ile Lys Trp Ile
                        55
     Cys Pro Thr Ala Pro Ser Gln Pro Ile Ser Leu Phe Gly Gly Phe Pro
                        70
15
     Ser Thr Ala Trp Phe Asp Val Val Asp Ile Asn Glu Asp Gly Pro Asp
                                       90
     Asp Met Glu Gly Leu Asp Val Ala Ala Ala His Val Ala Asn Leu Leu
                                   105
                100
     Ser Asn Glu Pro Ala Asp Ile Lys Leu Gly Val Gly Gly Phe Ser Met
20
                               120
     Gly Ala Ala Thr Ser Leu Tyr Ser Ala Thr Cys Phe Ala Leu Gly Lys
                           135
                                 140
     Tyr Gly Asn Gly Asn Pro Tyr Pro Ile Asn Leu Ser Ala Ile Ile Gly
                            155
                       150
25
     Leu Ser Gly Trp Leu Pro Cys Ala Lys Thr Leu Ala Gly Lys Leu Glu
                                      170 175
                   165
     Glu Glu Gln Ile Lys Asn Arg Ala Ala Ser Leu Pro Ile Val Val Cys
                                   185
                180
     His Gly Lys Ala Asp Asp Val Val Pro Phe Lys Phe Gly Glu Lys Ser
30
                               200
     Ser Gln Ala Leu Leu Ser Asn Gly Phe Lys Lys Val Thr Phe Lys Pro
                           215
                                              220
     Tyr Ser Ala Leu Gly His His Thr Ile Pro Gln Glu Leu Asp Glu Leu
           230
                                          235
35
     Cys Ala Trp Leu Thr Ser Thr Leu Ser Leu Glu Gly
                    245
     (2) INFORMATION FOR SEQ ID NO:45:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 155 amino acids
40
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
45
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..155
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011618
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:
     Met Glu Gly Leu Asp Val Ala Ala Ala His Val Ala Asn Leu Leu Ser
50
                                       10
     Asn Glu Pro Ala Asp Ile Lys Leu Gly Val Gly Gly Phe Ser Met Gly
                                   25
     Ala Ala Thr Ser Leu Tyr Ser Ala Thr Cys Phe Ala Leu Gly Lys Tyr
                               40
55
     Gly Asn Gly Asn Pro Tyr Pro Ile Asn Leu Ser Ala Ile Ile Gly Leu
     Ser Gly Trp Leu Pro Cys Ala Lys Thr Leu Ala Gly Lys Leu Glu Glu
     Glu Gln Ile Lys Asn Arg Ala Ala Ser Leu Pro Ile Val Val Cys His
60
                                      90
     Gly Lys Ala Asp Asp Val Val Pro Phe Lys Phe Gly Glu Lys Ser Ser
```

347 Gln Ala Leu Leu Ser Asn Gly Phe Lys Lys Val Thr Phe Lys Pro Tyr 115 120 Ser Ala Leu Gly His His Thr Ile Pro Gln Glu Leu Asp Glu Leu Cys 135 Ala Trp Leu Thr Ser Thr Leu Ser Leu Glu Gly 150 (2) INFORMATION FOR SEQ ID NO:46: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 643 base pairs 10 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 15 (A) NAME/KEY: -(B) LOCATION: 1..643 (D) OTHER INFORMATION: / Ceres Seq. ID 1011631 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46: agaatgtgat ggagatggga gtgaatgaga tgagtggtgg gattaaagaa gagaagaagg 20 agaaggctct tatgcttaga ttggactatg aatcagtcat ttccacttgg ggaggccaag ggatcccatg gaccgcccgg gtgccatctg aaatagacct cgacatggtt tgtttcccaa cccataccat gggtgaaagt ggagcagagg ctcatcatca caaccacttc cgcggcctag ggttacacct aggagatgct ggggatggag gaagagaggc tagggtttca agataccgag agaaaaggag gacaaggttg ttctccaaga agataaggta cgaggtacgt aaattgaatg cagagaaaag gcctcgcatg aaaggaaggt tcgtcaagag atcttcaatt ggtgttgctc 25 actaaagaac ttaattaatt atggatatta aattactttg ctctcatctt gcttttttgt tgctatagtt ttggtgattg ttagctttct ttttctgcat tcatagagaa ttttgcacgt ttttgtgagc tacgtatgta cataaatata tcaccaaaaa aatgtgacta tcttgtaagc actgatttat atagtcgata taacgtgaat tttgattgct ggc 30 (2) INFORMATION FOR SEQ ID NO:47: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 140 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 35 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..140 40 (D) OTHER INFORMATION: / Ceres Seq. ID 1011632 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47: Asn Val Met Glu Met Gly Val Asn Glu Met Ser Gly Gly Ile Lys Glu 1.5 10 Glu Lys Lys Glu Lys Ala Leu Met Leu Arg Leu Asp Tyr Glu Ser Val 45 25 Ile Ser Thr Trp Gly Gly Gln Gly Ile Pro Trp Thr Ala Arg Val Pro 40 Ser Glu Ile Asp Leu Asp Met Val Cys Phe Pro Thr His Thr Met Gly 55 50 Glu Ser Gly Ala Glu Ala His His His Asn His Phe Arg Gly Leu Gly 75 70 Leu His Leu Gly Asp Ala Gly Asp Gly Gly Arg Glu Ala Arg Val Ser 90 Arg Tyr Arg Glu Lys Arg Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg 55 105 100 110 Tyr Glu Val Arg Lys Leu Asn Ala Glu Lys Arg Pro Arg Met Lys Gly 115 120 Arg Phe Val Lys Arg Ser Ser Ile Gly Val Ala His 135 60 (2) INFORMATION FOR SEQ ID NO:48: (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 138 amino acids

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(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..138
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011633
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:
10
     Met Glu Met Gly Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys
                                          10
      Lys Glu Lys Ala Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser
                 20
                                      25
      Thr Trp Gly Gly Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu
15
                                  4.0
      Ile Asp Leu Asp Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser
                              5.5
      Gly Ala Glu Ala His His His Asn His Phe Arg Gly Leu Gly Leu His
                                              75
                         70
      Leu Gly Asp Ala Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr
20
                                          90
                     85
     Arg Glu Lys Arg Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu
                                     105
                 100
                                                          110
     Val Arg Lys Leu Asn Ala Glu Lys Arg Pro Arg Met Lys Gly Arg Phe
25
                                 120
     Val Lys Arg Ser Ser Ile Gly Val Ala His
         130
                             135
      (2) INFORMATION FOR SEQ ID NO:49:
          (i) SEQUENCE CHARACTERISTICS:
30
               (A) LENGTH: 136 amino acids
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..136
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011634
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:
40
     Met Gly Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu
              5
                                          10
                                                              15
      Lys Ala Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp
                                      25
      Gly Gly Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp
45
                                  40
      Leu Asp Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala
                              55
      Glu Ala His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly
                          70
                                              75
50
      Asp Ala Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu
                                          90
      Lys Arg Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg
                                      105
      Lys Leu Asn Ala Glu Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys
55
                                  120
      Arg Ser Ser Ile Gly Val Ala His
         130
                              135
      (2) INFORMATION FOR SEQ ID NO:50:
           (i) SEQUENCE CHARACTERISTICS:
60
                (A) LENGTH: 663 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
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(D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
 5
                (B) LOCATION: 1..663
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011714
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:
      gtgttcgata ataatgtcga caggagaagc gataccaaga gtcgccgtcg tcgttttcat
      teteaaegga aacteaatet tattaggteg eegeegttee teaateggea aeteeaettt
10
     cgctcttccc ggtggccacc tcgaattcgg agagagcttt gaagaatgtg cagcgagaga
      agtaatggag gaaacaggtc taaagattga aaagatgaag cttttgactg ttacaaacaa
                                                                            240
     tgtcttcaaa gaagcaccaa cgccatcaca ctacgtctct gtttcgatac gtgcggtgtt
                                                                            300
     ggtggatcca agtcaagaac cgaagaatat ggaaccagag aagtgtgaag gatgggattg
                                                                            360
     gtatgattgg gagaatctac caaagccttt gttttggcca cttgagaaat tgtttggaag
                                                                            420
15
     tggtttcaat cctttcactc atggtggtgg agactaatag atgtaagagt taatgattga
     tttgggattg aatgttgcaa attgggcatt tggtctagtg gtatgattct cgcttagggt
      gcgagaggtc ccgagttcaa ttctcggaat gcccctctct tttactcttt tttattgtat
                                                                            600
      ttttggtttc tgttttccta attagtaata aacttatcat gtgtcctatc gattttattt
     ctc
20
      (2) INFORMATION FOR SEQ ID NO:51:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 151 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..151
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011715
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:
      Cys Ser Ile Ile Met Ser Thr Gly Glu Ala Ile Pro Arg Val Ala Val
                                         10
      Val Val Phe Ile Leu Asn Gly Asn Ser Ile Leu Leu Gly Arg Arg Arg
35
                  20
                                      2.5
      Ser Ser Ile Gly Asn Ser Thr Phe Ala Leu Pro Gly Gly His Leu Glu
                                  40
      Phe Gly Glu Ser Phe Glu Glu Cys Ala Ala Arg Glu Val Met Glu Glu
                              55
40
      Thr Gly Leu Lys Ile Glu Lys Met Lys Leu Leu Thr Val Thr Asn Asn
                          70
                                              75
      Val Phe Lys Glu Ala Pro Thr Pro Ser His Tyr Val Ser Val Ser Ile
                                          90
      Arg Ala Val Leu Val Asp Pro Ser Gln Glu Pro Lys Asn Met Glu Pro
4.5
                  100
                                      105
      Glu Lys Cys Glu Gly Trp Asp Trp Tyr Asp Trp Glu Asn Leu Pro Lys
              115
                                  120
      Pro Leu Phe Trp Pro Leu Glu Lys Leu Phe Gly Ser Gly Phe Asn Pro
                              135
50
      Phe Thr His Gly Gly Asp
                          150
      (2) INFORMATION FOR SEQ ID NO:52:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 147 amino acids
55
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
60
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..147
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(D) OTHER INFORMATION: / Ceres Seq. ID 1011716

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:
     Met Ser Thr Gly Glu Ala Ile Pro Arg Val Ala Val Val Phe Ile
                                        1.0
     Leu Asn Gly Asn Ser Ile Leu Leu Gly Arg Arg Arg Ser Ser Ile Gly
 5
                                    25
                 20
     Asn Ser Thr Phe Ala Leu Pro Gly Gly His Leu Glu Phe Gly Glu Ser
                                40
     Phe Glu Glu Cys Ala Ala Arg Glu Val Met Glu Glu Thr Gly Leu Lys
                            55
10
     Ile Glu Lys Met Lys Leu Leu Thr Val Thr Asn Asn Val Phe Lys Glu
                                            75
     Ala Pro Thr Pro Ser His Tyı Val Ser Val Ser Ile Arg Ala Val Leu
                                        90
                     8.5
     Val Asp Pro Ser Gln Glu Pro Lys Asn Met Glu Pro Glu Lys Cys Glu
15
                                   105
                 100
     Gly Trp Asp Trp Tyr Asp Trp Glu Asn Leu Pro Lys Pro Leu Phe Trp
                                                   125
            115 120
     Pro Leu Glu Lys Leu Phe Gly Ser Gly Phe Asn Pro Phe Thr His Gly
        130 135
20
     Gly Gly Asp
     145
      (2) INFORMATION FOR SEQ ID NO:53:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 90 amino acids
25
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
30
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..90
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011717
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:
     Met Glu Glu Thr Gly Leu Lys Ile Glu Lys Met Lys Leu Leu Thr Val
35
              5
                                        10
      Thr Asn Asn Val Phe Lys Glu Ala Pro Thr Pro Ser His Tyr Val Ser
                 2.0
                                     25
                                                        30
      Val Ser Ile Arg Ala Val Leu Val Asp Pro Ser Gln Glu Pro Lys Asn
                                 40
40
      Met Glu Pro Glu Lys Cys Glu Gly Trp Asp Trp Tyr Asp Trp Glu Asn
                             55 .
                                             60
      Leu Pro Lys Pro Leu Phe Trp Pro Leu Glu Lys Leu Phe Gly Ser Gly
                         70
      Phe Asn Pro Phe Thr His Gly Gly Asp
45
                     85
      (2) INFORMATION FOR SEQ ID NO:54:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 524 base pairs
               (B) TYPE: nucleic acid
50
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
55
               (B) LOCATION: 1..524
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011784
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:
      aaaacagaac aaacacaaac acaaacagag aatcaacgaa gaagaaaaat ggtgaaggtg
                                                                          60
      atgtgggttt ccgttttagc tctggcggcg gcgattctcc ttttgacggt cccggtggca
                                                                         120
60
      gaaggggtga cgtgctcgcc tatgcagctg gcttcatgtg cggcggcgat gacgtcatct
                                                                         180
      tcgccgccat cggaggcgtg ttgcacaaag ctgagagagc agcagccatg cctttgtggg
                                                                         240
      tacatgagga accetaccet cegecaatac qttageteec ctaacgcaag gaaagtetee
                                                                         300
```

		351															
	acaa taat	igttt :aaat	.cg c	tgat gtgat	tata gato	ng to ga tt	cccaagctgt taaggaaatg tggttaatgc tggtcttatc ttgatgatta tcatcatcat tactttagaa atatgtttcg					ttcgtcagta ct gcttgaatgt ta				taaaa	360 420 480
5	(2)	<pre>INFORMATION FOR SEQ ID NO:55:    (i) SEQUENCE CHARACTERISTICS:         (A) LENGTH: 110 amino acids         (B) TYPE: amino acid         (C) STRANDEDNESS:</pre>															
10		(D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide															
15			( E	A) NA B) LC D) OI	CAT	ON:	11	10	: / (	 Ceres	Sec	ą. II	) 101	.1785	.· 5		
	Lys 1			QUENC Gln									Arg	Arg	Arg 15	Lys	
20		Val	Lys	Val 20		Trp	Val	Ser	Val 25		Ala	Leu	Ala	Ala 30		Ile	
	Leu	Leu	Leu 35	Thr	Val	Pro	Val	Ala 40	Glu	Gly	Val	Thr	Cys 45	Ser	Pro	Met	
	Gln	Leu 50	Ala	Ser	Cys	Ala	Ala 55	Ala	Met	Thr	Ser	Ser 60	Ser	Pro	Pro	Ser	
25	65			Cys		70					75					80	
				Asn	85					90					Asn 95	Ala	
30	(2)	INFO	ORMAT	Ser 100 TION QUENC A) LE	FOR CE CI	SEQ HARA(	ID N	0:50 STIC	105 6: CS:		ser	Pro	ser	110			
35		<pre>(B) TYPE: amino acid   (C) STRANDEDNESS:     (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE:</pre>															
40		/ w i '	(1	A) NA B) L( D) O' QUEN(	CAT:	ION: INF	1 DRMA	94 TION					D 10:	1178	6		
	Met 1			Val									Ala	Ala	Ala 15	Ile	
45	_	Leu	Leu	Thr 20	-	Pro	Val	Ala	Glu 25		Val	Thr	Cys	Ser 30		Met	
	Gln	Leu	Ala 35	Ser	Cys	Ala	Ala	Ala 40	Met	Thr	Ser	Ser	Ser 45	Pro	Pro	Ser	
50	Glu	Ala 50	Cys	Cys	Thr	Lys	Leu 55	Arg	Glu	Gln	Gln	Pro 60	Cys	Leu	Cys	Gly	
	65			Asn		70					75				Asn	Ala 80	
6.6				Ser	85					Pro 90	Ser	Pro	Ser	Cys			
55	(2)		) SE (. (	TION QUEN A) L B) T	CE C ENGT YPE:	HARA H: 9 ami	CTER 0 am no a	ISTI ino	CS:	s							
60			) MO (	C) S' D) T LECU ATUR	OPOL LE T	OGY:	lin					٠					

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(A) NAME/KEY: peptide
                (B) LOCATION: 1..90
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011787
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:
 5
     Asn Arg Thr Asn Thr Asn Thr Asn Arg Glu Ser Thr Lys Lys Lys Asn
                                      10
      Gly Glu Gly Asp Val Gly Phe Arg Phe Ser Ser Gly Gly Asp Ser
                                      25
      Pro Phe Asp Gly Pro Gly Gly Arg Arg Gly Asp Val Leu Ala Tyr Ala
10
                                  4.0
      Ala Gly Phe Met Cys Gly Gly Asp Asp Val Ile Phe Ala Ala Ile Gly
                              5.5
                                                  60
      Gly Val Leu His Lys Ala Glu Arg Ala Ala Ala Met Pro Leu Trp Val
                         70
                                             75
15
      His Glu Glu Pro Tyr Pro Pro Pro Ile Arg
                    85
      (2) INFORMATION FOR SEQ ID NO:58:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1007 base pairs
20
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
25
                (A) NAME/KEY: -
                (B) LOCATION: 1..1007
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011820
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:
      ccaaatttct ccatggatcg ttctttgaat ctcctcgatt tagccttagg gttcgatgag
                                                                             60
30
                                                                          120
      cagctagcta ttccatcgcc actaaatgga aaagtaatac taatagaaga ctgtgtagag
      acgagtggtt cctttgtact tcaccagcta atgaaacgtg ttctctcctc taactcctcc
      gacgcactta totttotogo tittgotogo cotttototo attatgatog aatottgogt
                                                                            240
      aaactgggat gtaatttagc tacccataag tcgaataatc gattggtgtt ctttgacatg
                                                                            300
      ctcatggtta agtgttcaga tggggatcaa atggaagaca atgtgagtgc agttgcgaaa
                                                                            360
35
      ctatttcggg agatacaaga aaccgttcga aagctacaga gtgtaacaag tggtaacata
                                                                            420
      actgttatgg tggatgacat gtctctgctg gaaattgcta ctaccggcag caactcagat
                                                                            480
      cacgtattgg acttcttgca ttattgccac acattaagtt ctgaaagcaa ttgttcattg
                                                                            540
      gtcatcctca atcatgaaga tatatacgcg agcatggaga gacctgcatt tttgctacag
                                                                            600
      atggtatgcc ttgcagatgt tgtgataaag gcagagcctt tagcctctgg tttagcaaat
                                                                            660
40
      gatgtacatg gccaattgac tgttctgaac aaagggataa gcaactcagg tagaggaagc
                                                                            720
      tcgaggaaca agttgcagaa ttttcaattc aggatcaagg aaaatggtat cgactatttc
                                                                            780
      tatcctggtt gcagaagctg aggattagac accgttctga gcattagcat ggtagtgaga
                                                                            840
      tgcatctaga cagttttgtt atttgcgttt aggttaaaca gagacactct gttcaacatt
                                                                            900
      tttactttag tgtctcttct ggttgtaact tgatgactta gctaaagttg aatgcttgag
                                                                            960
45
      acacagaaac ttgtagacat ggaaaattag taacattcta tatcttt
      (2) INFORMATION FOR SEQ ID NO:59:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 266 amino acids
                (B) TYPE: amino acid
50
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
55
                (B) LOCATION: 1..266
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011821
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:
      Pro Asn Phe Ser Met Asp Arg Ser Leu Asn Leu Leu Asp Leu Ala Leu
                    5
                                         10
60
      Gly Phe Asp Glu Gln Leu Ala Ile Pro Ser Pro Leu Asn Gly Lys Val
```

20 25 30

Ile Leu Ile Glu Asp Cys Val Glu Thr Ser Gly Ser Phe Val Leu His

```
Gln Leu Met Lys Arg Val Leu Ser Ser Asn Ser Ser Asp Ala Leu Ile
     Phe Leu Ala Phe Ala Arg Pro Phe Ser His Tyr Asp Arg Ile Leu Arg
 5
                       70
                                         75
     Lys Leu Gly Cys Asn Leu Ala Thr His Lys Ser Asn Asn Arg Leu Val
                                     90
     Phe Phe Asp Met Leu Met Val Lys Cys Ser Asp Gly Asp Gln Met Glu
             100 105
10
     Asp Asn Val Ser Ala Val Ala Lys Leu Phe Arg Glu Ile Gln Glu Thr
           115 120
                                     125
     Val Arg Lys Leu Gln Ser Val Thr Ser Gly Asn Ile Thr Val Met Val
                               140
       130 135
     Asp Asp Met Ser Leu Leu Glu Ile Ala Thr Thr Gly Ser Asn Ser Asp
                            155 160
15
          150
     His Val Leu Asp Phe Leu His Tyr Cys His Thr Leu Ser Ser Glu Ser
                 165 170 175
     Asn Cys Ser Leu Val Ile Leu Asn His Glu Asp Ile Tyr Ala Ser Met
               180
                     185 190
20
     Glu Arg Pro Ala Phe Leu Leu Gln Met Val Cys Leu Ala Asp Val Val
            195 200
     Ile Lys Ala Glu Pro Leu Ala Ser Gly Leu Ala Asn Asp Val His Gly
             215
     Gln Leu Thr Val Leu Asn Lys Gly Ile Ser Asn Ser Gly Arg Gly Ser
25
                      230
                                         235
     Ser Arg Asn Lys Leu Gln Asn Phe Gln Phe Arg Ile Lys Glu Asn Gly
     Ile Asp Tyr Phe Tyr Pro Gly Cys Arg Ser
                260
30
     (2) INFORMATION FOR SEQ ID NO:60:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 262 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
35
              (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..262
40
              (D) OTHER INFORMATION: / Ceres Seq. ID 1011822
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:
     Met Asp Arg Ser Leu Asn Leu Leu Asp Leu Ala Leu Gly Phe Asp Glu
         5
                                     10
     Gln Leu Ala Ile Pro Ser Pro Leu Asn Gly Lys Val Ile Leu Ile Glu
45
     Asp Cys Val Glu Thr Ser Gly Ser Phe Val Leu His Gln Leu Met Lys
                              40
     Arg Val Leu Ser Ser Asn Ser Ser Asp Ala Leu Ile Phe Leu Ala Phe
50
     Ala Arg Pro Phe Ser His Tyr Asp Arg Ile Leu Arg Lys Leu Gly Cys
     Asn Leu Ala Thr His Lys Ser Asn Asn Arg Leu Val Phe Phe Asp Met
     Leu Met Val Lys Cys Ser Asp Gly Asp Gln Met Glu Asp Asn Val Ser
55
                                 105
                100
     Ala Val Ala Lys Leu Phe Arg Glu Ile Gln Glu Thr Val Arg Lys Leu
                             120
     Gln Ser Val Thr Ser Gly Asn Ile Thr Val Met Val Asp Asp Met Ser
                       135
60
     Leu Leu Glu Ile Ala Thr Thr Gly Ser Asn Ser Asp His Val Leu Asp
                       150
                                        155
     Phe Leu His Tyr Cys His Thr Leu Ser Ser Glu Ser Asn Cys Ser Leu
```

										354						
	Val	Ile	Leu		165 His	Glu	Asp	Ile	Tyr	170 Ala	Ser	Met	Glu		175 Pro	Ala
5	Phe	Leu		180 Gln	Met	Val	Cys		185 Ala	Asp	Val	Val		190 Lys	Ala	Glu
J	Pro	Leu 210	195 Ala	Ser	Gly	Leu	Ala 215	200 Asn	Asp	Val	His	Gly 220	205 Gln	Leu	Thr	Val
	Leu 225		Lys	Gly	Ile	Ser 230		Ser	Gly	Arg	Gly 235		Ser	Arg	Asn	Lys 240
10		Gln	Asn	Phe	Gln 245		Arg	Ile	Lys	Glu 250		Gly	Ile	Asp	Tyr 255	
	Tyr	Pro	Gly	Cys 260	Arg	Ser										
15	<ul> <li>(2) INFORMATION FOR SEQ ID NO:61:</li> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 216 amino acids</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS:</li> <li>(D) TOPOLOGY: linear</li> </ul>															
20	(ii) MOLECULE TYPE: peptide (ix) FEATURE:  (A) NAME/KEY: peptide  (B) LOCATION: 1216  (D) OTHER INFORMATION: / Ceres Seq. ID 1011823															
25	Met		SEÇ	QUENC	CE DI	ESCR	IPTI	: NC	SEQ :	ID NO	61:61	:				Leu
	1 Ala	Phe	Ala		5 Pro	Phe	Ser	His	Tyr	10 Asp	Arg	Ile	Leu		15 Lys	Leu
30	Gly	Cys	Asn 35	20 Leu	Ala	Thr	His	Lys 40	25 Ser	Asn	Asn	Arg	Leu 45	30 Val	Phe	Phe
	Asp	Met 50		Met	Val	Lys	Cys 55		Asp	Gly	Asp	Gln 60		Glu	Asp	Asn
35	65					70			Arg		75					80
					85				Asn	90					95	
4.0				100					Thr 105	_				110		
40			115					120	Thr				125			Cys
		130					135		Cys		-	140				_
45	145					150			Ala		155					160
					165				Asn	170					175	
50	Asn	Lys		180 Gln	Asn	Phe	Gln		185 Arg	Ile	Lys	Glu		_		Asp
	Tyr		195 Tyr	Pro	Gly	Суѕ	_	200 Ser					205			
55	210 215 (2) INFORMATION FOR SEQ ID NO:62: (3) (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 584 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single															
60			) MO ) FE	LECU ATUR	LE T	YPE:			nomi	c)						

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(B) LOCATION: 1..584
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011874
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:
      tcaattaaca taaccttcgc cgtcgatgaa gctttccctt caattcccac taccttcagc
                                                                              60
5
      gtogoaacga aacaacacta ogacqtaaaa coaatocaco atgaaqttta tgaoggagaa
                                                                             120
      aggaagatat acgacatcag ccaccagtac acgccggagt tgccggtttg ggagtcttca
                                                                             180
      gaaggactag ggaactttct tagacttgcc gtgagtatga agaatggatc cgatgctaat
                                                                             240
      atctcgaaga tggaactatc tgttcactct ggaactcatg ttgatgcacc aggccatttc
                                                                             300
      catgaccatt attatgagtc tggttttgat actgattcac ttgatcttca aatccttaat
                                                                             360
10
      ggtaaacatt tttggaactc ttttttttt tttttttaa gtagctagtg atcgtttttg
                                                                             420
      aaattggttt ctgtgaattt aatttatagg tcctgcttta ttggttgatg ttccaagaga
                                                                             480
      taagaacatt tcaggtttgt tgagagtttt ctaacattga aatttttaca aaatcttcat
                                                                             540
      taaatgtttg ttacaaaatt tttgtgtaat gatttggttg ttgt
      (2) INFORMATION FOR SEQ ID NO:63:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 135 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..135
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011875
25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:
      Ser Ile Asn Ile Thr Phe Ala Val Asp Glu Ala Phe Pro Ser Ile Pro
                                          10
      Thr Thr Phe Ser Val Ala Thr Lys Gln His Tyr Asp Val Lys Pro Ile
                                      25
30
      His His Glu Val Tyr Asp Gly Glu Arg Lys Ile Tyr Asp Ile Ser His
                                  4 0
      Gln Tyr Thr Pro Glu Leu Pro Val Trp Glu Ser Ser Glu Gly Leu Gly
                              55
      Asn Phe Leu Arg Leu Ala Val Ser Met Lys Asn Gly Ser Asp Ala Asn
35
                          70
                                              7.5
      Ile Ser Lys Met Glu Leu Ser Val His Ser Gly Thr His Val Asp Ala
                                           90
                      85
      Pro Gly His Phe His Asp His Tyr Tyr Glu Ser Gly Phe Asp Thr Asp
                                      105
40
      Ser Leu Asp Leu Gln Ile Leu Asn Gly Lys His Phe Trp Asn Ser Phe
              115
                                  120
      Phe Phe Phe Leu Ser Ser
          130
                              135
      (2) INFORMATION FOR SEQ ID NO:64:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 626 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..626
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011981
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:
      acatectaat egaaaaaac eetageaaaa acetetetet eteteteaat teaettetee
                                                                              60
      tectacaatg gettggegea acgeaggate tgetgetegt tetttegtet eegecacege
                                                                             120
      aagatcacca tototoogtt otoccaccac ggogottoot ogcotoogto otoctcaato
                                                                             180
      ctccttacct agccgtcgct tcaccttctc gtcaccttcc aggaatctag gagccctagg
                                                                             240
60
                                                                             300
      ttgcacacag tcgttcttgc ctctgtacag tgttgtggct acttctcaac tgacatctca
      ccttaacgtt aatttgcgag ctttctgcga gctgtctaac ggtacttgaa gaggcaaaag
                                                                             360
      agtgtaattt cactgcgcat gcgtgaatca gtactttgaa gggatgtgag atcgaaattg
                                                                             420
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tgggattacc tcagcaaaaa cttgcttaat gtgttgcctg ttttagaagt tttagtttca
                                                                             480
     caatgtgtgg atttattagt tgtagagctt ttcgtatgct tggcacaatg gttttttctc
                                                                             540
     agagaacctt tttttgtttt ggagaaatgc atagcagcat taataagtgt attggatcag
                                                                             600
     aactgaaaca atgttgttac tgattc
 5
     (2) INFORMATION FOR SEQ ID NO:65:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 93 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
10
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..93
15
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011982
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:
      Met Ala Trp Arg Asn Ala Gly Ser Ala Ala Arg Ser Phe Val Ser Ala
                                          10
      Thr Ala Arg Ser Pro Ser Leu Arg Ser Pro Thr Thr Ala Leu Pro Arg
20
                  2.0
                                      25
      Leu Arg Pro Pro Gln Ser Ser Leu Pro Ser Arg Arg Phe Thr Phe Ser
                                  40
      Ser Pro Ser Arg Asn Leu Gly Ala Leu Gly Cys Thr Gln Ser Phe Leu
25
      Pro Leu Tyr Ser Val Val Ala Thr Ser Gln Leu Thr Ser His Leu Asn
                          70
      Val Asn Leu Arg Ala Phe Cys Glu Leu Ser Asn Gly Thr
                      85
      (2) INFORMATION FOR SEQ ID NO:66:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 75 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011983
40
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:
      Ile Leu Ile Glu Lys Asn Pro Ser Lys Asn Leu Ser Leu Ser Leu Asn
                                          10
      Ser Leu Leu Leu Gln Trp Leu Gly Ala Thr Gln Asp Leu Leu
                  20
                                      25
45
      Val Leu Ser Ser Pro Pro Pro Gln Asp His His Leu Ser Val Leu Pro
                                  40
      Pro Arg Arg Phe Leu Ala Ser Val Leu Leu Asn Pro Pro Tyr Leu Ala
                              55
      Val Ala Ser Pro Ser Arg His Leu Pro Gly Ile
50
                          70
      (2) INFORMATION FOR SEQ ID NO:67:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 46 amino acids
                 (B) TYPE: amino acid
55
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
60
                 (B) LOCATION: 1..46
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1011984
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:
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357
     Met Cys Gly Phe Ile Ser Cys Arg Ala Phe Arg Met Leu Gly Thr Met
                                          10
     Val Phe Ser Gln Arg Thr Phe Phe Cys Phe Gly Glu Met His Ser Ser
                                      25
5
     Ile Asn Lys Cys Ile Gly Ser Glu Leu Lys Gln Cys Cys Tyr
             35
                                 40
      (2) INFORMATION FOR SEQ ID NO:68:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 468 base pairs
10
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
15
                (A) NAME/KEY: -
                (B) LOCATION: 1..468
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014547
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
      atagaatata accatattcc ggacatgaag atcgtgacat tggtactcgt cgtcttcgtc
                                                                              60
20
      atactttcga catcattccc ggctgccatc aaagccgaag acacgggaga tacaggaaat
                                                                             120
      gtgggagtga catgtgacgc aaggcagctt cagccttgcc ttgccgcgat tacgggagga
                                                                             180
      ggacaaccct cgggtqcatg ttgtqcaaag cttacagagc aacagtcatg cctatgtggt
                                                                             240
      ttcgctaaga accetgcgtt cgcacagtac attagctctc caaacgctcg caaagtgctc
                                                                             300
      cttgcttgca atgttgctta tcccacttgt tgaaacttat ctagatttta taaataaata
                                                                             360
25
      aacgaaagga aataaattac attatatcaa acgttatgat acaattcaac cgtttgtgtt
                                                                            420
      aatgtactgg cttacatggt taaataaagt ttaatttctt ggttgctc
      (2) INFORMATION FOR SEQ ID NO:69:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 110 amino acids
30
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..110
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014548
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:
      Ile Glu Tyr Asn His Ile Pro Asp Met Lys Ile Val Thr Leu Val Leu
40
                      5
                                           10
      Val Val Phe Val Ile Leu Ser Thr Ser Phe Pro Ala Ala Ile Lys Ala
                  20
                                      25
      Glu Asp Thr Gly Asp Thr Gly Asn Val Gly Val Thr Cys Asp Ala Arg
                                                       45
                                  40
45
      Gln Leu Gln Pro Cys Leu Ala Ala Ile Thr Gly Gly Gln Pro Ser
                              55
                                                   60
      Gly Ala Cys Cys Ala Lys Leu Thr Glu Gln Gln Ser Cys Leu Cys Gly
                          70
                                               75
      Phe Ala Lys Asn Pro Ala Phe Ala Gln Tyr Ile Ser Ser Pro Asn Ala
50
                                          90
                     85
      Arg Lys Val Leu Leu Ala Cys Asn Val Ala Tyr Pro Thr Cys
                  100
                                       105
      (2) INFORMATION FOR SEQ ID NO:70:
           (i) SEQUENCE CHARACTERISTICS:
55
                 (A) LENGTH: 102 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
60
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..102
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(D) OTHER INFORMATION: / Ceres Seq. ID 1014549
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:
     Met Lys Ile Val Thr Leu Val Leu Val Val Phe Val Ile Leu Ser Thr
                                          10
                      5
5
     Ser Phe Pro Ala Ala Ile Lys Ala Glu Asp Thr Gly Asp Thr Gly Asn
                  20
                                      25
     Val Gly Val Thr Cys Asp Ala Arg Gln Leu Gln Pro Cys Leu Ala Ala
      Ile Thr Gly Gly Gly Gln Pro Ser Gly Ala Cys Cys Ala Lys Leu Thr
10
      Glu Gln Gln Ser Cys Leu Cys Gly Phe Ala Lys Asn Pro Ala Phe Ala
                                              75
      Gln Tyr Ile Ser Ser Pro Asn Ala Arg Lys Val Leu Leu Ala Cys Asn
                                          90
                      85
15
      Val Ala Tyr Pro Thr Cys
                  100
      (2) INFORMATION FOR SEQ ID NO:71:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1303 base pairs
20
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
25
                (A) NAME/KEY: -
                (B) LOCATION: 1..1303
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014995
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:
      aaaactcaag tttcttgcaa ttacctactt acaacaaagc aactcgactc gaacaaacac
                                                                              60
30
      atccatggct ccttctgcgc aacctcttcc tgtgagtgtt tcggatgaaa aatatgcgaa
                                                                             120
      tgtcaagtgg gaagagttgg cattcaagtt tgttcgtacg gattatatgt atgttgcgaa
                                                                             180
      gtgcaatcat ggagagagtt ttcaagaggg gaagattctt ccttttgctg atttgcaact
                                                                             240
      taacccttgc gctgctgttc ttcagtatgg ccagggttta tatgaaggac tgaaagctta
                                                                             300
      caggacagaa gatggtcgga ttctgctatt ccgaccagac caaaacggtc tccgccttca
                                                                             360
35
      agcoggaget gacagactet atatgeetta teetteggte gateaatteg teteegeeat
                                                                             420
      caaacaagtt gctcttgcca acaagaaatg gattcctcct ccggggaaag gaacattgta
                                                                             480
      tattaggcct atcttgtttg ggagtggtcc gattcttggt tcatttccca ttcctgagac
                                                                             540
      caccttcaca gcttttgcct gtcctgttgg acgttatcat aaggataact ctggtttgaa
                                                                             600
      tctgaaaatc gaagatcagt ttcgtcgagc ttttcctagt ggaactggtg gtgtgaagag
                                                                             660
40
                                                                             720
      catcacaaac tattgtcctg tttggatacc attggcagag gcgaaaaaac aaggtttctc
      tgatattttg tttttggatg ctgcaactgg caaaaacatt gaagaacttt tcgcagctaa
                                                                             780
                                                                             840
      tgtttttatg ctcaagggca atgttgtatc gacaccaaca attgcaggaa ctattttgcc
                                                                             900
      cggagtcact cgaaactgcg taatggaatt gtgtcgtgat ttcggctacc aggtcgagga
                                                                             960
      acgtacgatt cctctaqtgg actttctcga tgcggacgaa gctttctgta ctggcactgc
45
      ttccattgtg actagtattg catccgtaac ctttaaagac aaaaagaccg gattcaaaac
                                                                            1020
                                                                            1080
      agggaagaaa cattggctgc gaagctatac gagacgttaa gtgatatcca gacgggtcgg
      gtcgaggata ccaagggatg gacggtggag attgaccgcc agggctgaaa gttgaaactg
                                                                            1140
      taacttgatg ctaaatatgt gtgtgtgtgt atacaaaact tatgtaagaa acatctgaag
                                                                            1200
      atgtetetga tetttgtgat tgtgatgate atgeatgtee tategttgeg atgtatttta
                                                                             1260
50
      taaatgttcg tctgtaagtt atttaataac tatggctttt tgc
      (2) INFORMATION FOR SEQ ID NO:72:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 352 amino acids
                 (B) TYPE: amino acid
55
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
60
                 (B) LOCATION: 1..352
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1014996
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:
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359 Lys Leu Lys Phe Leu Ala Ile Thr Tyr Leu Gln Gln Ser Asn Ser Thr 10 Arg Thr Asn Thr Ser Met Ala Pro Ser Ala Gln Pro Leu Pro Val Ser 25 5 Val Ser Asp Glu Lys Tyr Ala Asn Val Lys Trp Glu Glu Leu Ala Phe 4.0 Lys Phe Val Arg Thr Asp Tyr Met Tyr Val Ala Lys Cys Asn His Gly 55 Glu Ser Phe Gln Glu Gly Lys Ile Leu Pro Phe Ala Asp Leu Gln Leu 10 70 75 Asn Pro Cys Ala Ala Val Leu Gln Tyr Gly Gln Gly Leu Tyr Glu Gly 90 Leu Lys Ala Tyr Arg Thr Glu Asp Gly Arg Ile Leu Leu Phe Arg Pro 100 105 Asp Gln Asn Gly Leu Arg Leu Gln Ala Gly Ala Asp Arg Leu Tyr Met 15 120 Pro Tyr Pro Ser Val Asp Gln Phe Val Ser Ala Ile Lys Gln Val Ala 135 Leu Ala Asn Lys Lys Trp Ile Pro Pro Pro Gly Lys Gly Thr Leu Tyr 20 150 155 Ile Arg Pro Ile Leu Phe Gly Ser Gly Pro Ile Leu Gly Ser Phe Pro 165 170 Ile Pro Glu Thr Thr Phe Thr Ala Phe Ala Cys Pro Val Gly Arg Tyr 185 25 His Lys Asp Asn Ser Gly Leu Asn Leu Lys Ile Glu Asp Gln Phe Arg 200 Arg Ala Phe Pro Ser Gly Thr Gly Gly Val Lys Ser Ile Thr Asn Tyr 215 220 Cys Pro Val Trp Ile Pro Leu Ala Glu Ala Lys Lys Gln Gly Phe Ser 230 30 235 Asp Ile Leu Phe Leu Asp Ala Ala Thr Gly Lys Asn Ile Glu Glu Leu 245 250 255 Phe Ala Ala Asn Val Phe Met Leu Lys Gly Asn Val Val Ser Thr Pro 265 35 Thr Ile Ala Gly Thr Ile Leu Pro Gly Val Thr Arg Asn Cys Val Met 280 Glu Leu Cys Arg Asp Phe Gly Tyr Gln Val Glu Glu Arg Thr Ile Pro 290 295 300 Leu Val Asp Phe Leu Asp Ala Asp Glu Ala Phe Cys Thr Gly Thr Ala 40 315 310 Ser Ile Val Thr Ser Ile Ala Ser Val Thr Phe Lys Asp Lys Lys Thr 325 330 Gly Phe Lys Thr Gly Lys Lys His Trp Leu Arg Ser Tyr Thr Arg Arg 340 45 (2) INFORMATION FOR SEQ ID NO:73: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 331 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..331 55 (D) OTHER INFORMATION: / Ceres Seq. ID 1014997 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:73: Met Ala Pro Ser Ala Gln Pro Leu Pro Val Ser Val Ser Asp Glu Lys 10 Tyr Ala Asn Val Lys Trp Glu Glu Leu Ala Phe Lys Phe Val Arg Thr 60 25 20 Asp Tyr Met Tyr Val Ala Lys Cys Asn His Gly Glu Ser Phe Gln Glu

360 Gly Lys Ile Leu Pro Phe Ala Asp Leu Gln Leu Asn Pro Cys Ala Ala 55 Val Leu Gln Tyr Gly Gln Gly Leu Tyr Glu Gly Leu Lys Ala Tyr Arg 75 70 5 Thr Glu Asp Gly Arg Ile Leu Leu Phe Arg Pro Asp Gln Asn Gly Leu 85 90 Arg Leu Gln Ala Gly Ala Asp Arg Leu Tyr Met Pro Tyr Pro Ser Val 105 Asp Gln Phe Val Ser Ala Ile Lys Gln Val Ala Leu Ala Asn Lys Lys 10 120 Trp Ile Pro Pro Pro Gly Lys Gly Thr Leu Tyr Ile Arg Pro Ile Leu 135 Phe Gly Ser Gly Pro Ile Leu Gly Ser Phe Pro Ile Pro Glu Thr Thr 150 155 Phe Thr Ala Phe Ala Cys Pro Val Gly Arg Tyr His Lys Asp Asn Ser 15 170 165 Gly Leu Asn Leu Lys Ile Glu Asp Gln Phe Arg Arg Ala Phe Pro Ser 185 Gly Thr Gly Gly Val Lys Ser Ile Thr Asn Tyr Cys Pro Val Trp Ile 20 200 Pro Leu Ala Glu Ala Lys Lys Gln Gly Phe Ser Asp Ile Leu Phe Leu 215 220 Asp Ala Ala Thr Gly Lys Asn Ile Glu Glu Leu Phe Ala Ala Asn Val 230 235 25 Phe Met Leu Lys Gly Asn Val Val Ser Thr Pro Thr Ile Ala Gly Thr 245 250 Ile Leu Pro Gly Val Thr Arg Asn Cys Val Met Glu Leu Cys Arg Asp 265 Phe Gly Tyr Gln Val Glu Glu Arg Thr Ile Pro Leu Val Asp Phe Leu 30 280 Asp Ala Asp Glu Ala Phe Cys Thr Gly Thr Ala Ser Ile Val Thr Ser 295 Ile Ala Ser Val Thr Phe Lys Asp Lys Lys Thr Gly Phe Lys Thr Gly 305 310 315 35 Lys Lys His Trp Leu Arg Ser Tyr Thr Arg Arg 325 (2) INFORMATION FOR SEQ ID NO:74: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 297 amino acids 40 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 45 (A) NAME/KEY: peptide (B) LOCATION: 1..297 (D) OTHER INFORMATION: / Ceres Seq. ID 1014998 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:74: Met Tyr Val Ala Lys Cys Asn His Gly Glu Ser Phe Gln Glu Gly Lys 50 10 Ile Leu Pro Phe Ala Asp Leu Gln Leu Asn Pro Cys Ala Ala Val Leu 25 Gln Tyr Gly Gln Gly Leu Tyr Glu Gly Leu Lys Ala Tyr Arg Thr Glu 55 Asp Gly Arg Ile Leu Leu Phe Arg Pro Asp Gln Asn Gly Leu Arg Leu Gln Ala Gly Ala Asp Arg Leu Tyr Met Pro Tyr Pro Ser Val Asp Gln Phe Val Ser Ala Ile Lys Gln Val Ala Leu Ala Asn Lys Lys Trp Ile 60 90 Pro Pro Pro Gly Lys Gly Thr Leu Tyr Ile Arg Pro Ile Leu Phe Gly

361 Ser Gly Pro Ile Leu Gly Ser Phe Pro Ile Pro Glu Thr Thr Phe Thr 115 120 Ala Phe Ala Cys Pro Val Gly Arg Tyr His Lys Asp Asn Ser Gly Leu 135 140 5 Asn Leu Lys Ile Glu Asp Gln Phe Arg Arg Ala Phe Pro Ser Gly Thr 155 Gly Gly Val Lys Ser Ile Thr Asn Tyr Cys Pro Val Trp Ile Pro Leu 170 Ala Glu Ala Lys Lys Gln Gly Phe Ser Asp Ile Leu Phe Leu Asp Ala 10 180 185 Ala Thr Gly Lys Asn Ile Glu Glu Leu Phe Ala Ala Asn Val Phe Met 205 200 Leu Lys Gly Asn Val Val Ser Thr Pro Thr Ile Ala Gly Thr Ile Leu 210 215 15 Pro Gly Val Thr Arg Asn Cys Val Met Glu Leu Cys Arg Asp Phe Gly 230 235 Tyr Gln Val Glu Glu Arg Thr Ile Pro Leu Val Asp Phe Leu Asp Ala 245 250 Asp Glu Ala Phe Cys Thr Gly Thr Ala Ser Ile Val Thr Ser Ile Ala 20 265 Ser Val Thr Phe Lys Asp Lys Lys Thr Gly Phe Lys Thr Gly Lys Lys 280 His Trp Leu Arg Ser Tyr Thr Arg Arg 295 25 (2) INFORMATION FOR SEQ ID NO:75: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 789 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..789 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1015323 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:75: gggttaggtt tgtcttcaaa atcgtcagaa gaagaagaag aagtcaaaga cgataacaat 60 gacgtcatca tcgtacctcc gatttgcgat cgccgtcgta gctttcctct caatcaccac 120 180 catcaccacc gcacgtcctt gcaaaacctt cctaatctcc tcatattccc tctcaataac 40 240 cccagaaaac cctaacctcg aatccgattt cacctccaca cgattcgtca ccgttttcac 300 aatccgtcgt ctaaatcctc accacqtqqt tcctttcttc qtcaatcgtc gtcacqaqaa 360 gccacagatc caateggate gttccctccc tttgatctct gataacatca attcgtttag agategaace agagatatte ttagegtegt egtegegett ttgtteggtg ttggttgtgg 420 tgctttaacc gctgctacta tgtatcttgt ttgggctctt gttgtgaatc gtcaaagcta 480 45 tgattttgag gaagaagaag atgattatga qaatgatgaa tctgatgctg ctagcttgaa 540 gaaattgggt tatgttaaga ttccagctcc agctcctgct cctgtgaagg aagctgcttg 600 atttgaggtt gtgaatttca gtttctgcat attatcatga tttacgttgt tgcttatcta 660 tgttgttgcg tacaagtaaa ttcgtctctg ttatggattt atctaaactt actatactct 720 aagtatcagt gtttttacga tttgtagttg atgtttctta tatatata taagtgtgat 780 50 tatgatgtc (2) INFORMATION FOR SEQ ID NO:76: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 180 amino acids (B) TYPE: amino acid 55 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 60 (B) LOCATION: 1..180

(D) OTHER INFORMATION: / Ceres Seq. ID 1015324

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:76:

362 Met Thr Ser Ser Tyr Leu Arg Phe Ala Ile Ala Val Val Ala Phe 10 Leu Ser Ile Thr Thr Ile Thr Thr Ala Arg Pro Cys Lys Thr Phe Leu 25 Ile Ser Ser Tyr Ser Leu Ser Ile Thr Pro Glu Asn Pro Asn Leu Glu 40 Ser Asp Phe Thr Ser Thr Arg Phe Val Thr Val Phe Thr Ile Arg Arg 55 Leu Asn Pro His His Val Val Pro Phe Phe Val Asn Arg Arg His Glu 10 70 Lys Pro Gln Ile Gln Ser Asp Arg Ser Leu Pro Leu Ile Ser Asp Asn 85 Ile Asn Ser Phe Arg Asp Arg Thr Arg Asp Ile Leu Ser Val Val 105 15 Ala Leu Leu Phe Gly Val Gly Cys Gly Ala Leu Thr Ala Ala Thr Met 120 Tyr Leu Val Trp Ala Leu Val Val Asn Arg Gln Ser Tyr Asp Phe Glu 135 Glu Glu Glu Asp Asp Tyr Glu Asn Asp Glu Ser Asp Ala Ala Ser Leu 20 150 155 Lys Lys Leu Gly Tyr Val Lys Ile Pro Ala Pro Ala Pro Ala Pro Val 170 165 Lys Glu Ala Ala 25 (2) INFORMATION FOR SEQ ID NO:77: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 495 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..495 (D) OTHER INFORMATION: / Ceres Seq. ID 1016486 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77: acttecteat tageatteta cagtetgeae acttateaet tetttegaaa teteteteta 60 tccctctcaa aaatgaagct ctctgtgcgt tttatctccg ctgctcttct cttgttcatg 120 gtattcattg ccacagggat gggtccagtc accgtggagg cacgcacgtg tgagtcaaag 180 40 agccataggt tcaagggtcc atgtgtgagc acacacaact gtgcaaacgt gtgccacaac 240 gaaggetteg geggaggtaa atgeegtgga tteegtegte gttgetaetg cacaagacae 300 tgctgatcca tccattctca tgactcaaat cttcgatcca tcgtcagtgt gttacttctt 360 tottatotaa atottoogta oggtacoata togtacogta catgagtgtt ttotcaaata 420 agtcqttqqt ttqtqtttt ccqqttttaa tqtaatgtta aatcaattaa tggcttttaa 480 45 tatattgtat tatgg (2) INFORMATION FOR SEQ ID NO:78: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 101 amino acids (B) TYPE: amino acid 50 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 55 (B) LOCATION: 1..101 (D) OTHER INFORMATION: / Ceres Seq. ID 1016487 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:78: Thr Ser Ser Leu Ala Phe Tyr Ser Leu His Thr Tyr His Phe Phe Arg 10 60 Asn Leu Ser Leu Ser Leu Ser Lys Met Lys Leu Ser Val Arg Phe Ile

20 25 30 Ser Ala Ala Leu Leu Phe Met Val Phe Ile Ala Thr Gly Met Gly

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40
                                                    4.5
     Pro Val Thr Val Glu Ala Arg Thr Cys Glu Ser Lys Ser His Arg Phe
                            55
     Lys Gly Pro Cys Val Ser Thr His Asn Cys Ala Asn Val Cys His Asn
 5
                        70
                                    75
     Glu Gly Phe Gly Gly Gly Lys Cys Arg Gly Phe Arg Arg Cys Tyr
                    85
                                        90
     Cys Thr Arg His Cys
                 100
10
      (2) INFORMATION FOR SEQ ID NO:79:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 77 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
15
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..77
20
               (D) OTHER INFORMATION: / Ceres Seq. ID 1016488
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:
     Met Lys Leu Ser Val Arg Phe Ile Ser Ala Ala Leu Leu Leu Phe Met
              5
                                     10
      Val Phe Ile Ala Thr Gly Met Gly Pro Val Thr Val Glu Ala Arg Thr
25
                 20
                                     25
      Cys Glu Ser Lys Ser His Arg Phe Lys Gly Pro Cys Val Ser Thr His
             35
                                 4.0
      Asn Cys Ala Asn Val Cys His Asn Glu Gly Phe Gly Gly Lys Cys
                      55
30
      Arg Gly Phe Arg Arg Cys Tyr Cys Thr Arg His Cys
                         70
      (2) INFORMATION FOR SEQ ID NO:80:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 62 amino acids
35
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..62
               (D) OTHER INFORMATION: / Ceres Seq. ID 1016489
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:
      Met Val Phe Ile Ala Thr Gly Met Gly Pro Val Thr Val Glu Ala Arg
45
                                      10
      Thr Cys Glu Ser Lys Ser His Arg Phe Lys Gly Pro Cys Val Ser Thr
                                     25
      His Asn Cys Ala Asn Val Cys His Asn Glu Gly Phe Gly Gly Lys
                          40
50
      Cys Arg Gly Phe Arg Arg Cys Tyr Cys Thr Arg His Cys
                            55
      (2) INFORMATION FOR SEQ ID NO:81:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 325 base pairs
55
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
60
                (A) NAME/KEY: -
                (B) LOCATION: 1..325
                (D) OTHER INFORMATION: / Ceres Seq. ID 1018341
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:81:
      atcatcaaac acaaaacaat caatcaaaaa cataaaattc aaagtgtaag caaaaccaga
      acgagagaaa atggacaaca agcaaaacgc gagctaccaa gccggtcaag ccactggcca
      gactaaggag aaggeeggtg gaatgatgga caaggeeaag gatgetgetg etteagetea
                                                                            180
      agactccttg caacagactg gacaacaaat gaaggagaag gcgcaaggag ctgctgatgt
                                                                            240
     cgtcaaggac aagaccggca tgaacaaaag ccactaagca agcatcttgg acttcttttc
                                                                            300
      taattaattc ccctcttcga cttgt
      (2) INFORMATION FOR SEQ ID NO:82:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 68 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
15
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..68
                (D) OTHER INFORMATION: / Ceres Seq. ID 1018342
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:82:
20
      Met Asp Asn Lys Gln Asn Ala Ser Tyr Gln Ala Gly Gln Ala Thr Gly
                                          10
      Gln Thr Lys Glu Lys Ala Gly Gly Met Met Asp Lys Ala Lys Asp Ala
                                      25
      Ala Ser Ala Gln Asp Ser Leu Gln Gln Thr Gly Gln Gln Met Lys
25
                                  40
      Glu Lys Ala Gln Gly Ala Ala Asp Val Val Lys Asp Lys Thr Gly Met
      Asn Lys Ser His
      65
30
      (2) INFORMATION FOR SEQ ID NO:83:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 44 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
35
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..44
40
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1018343
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:83:
      Met Met Asp Lys Ala Lys Asp Ala Ala Ala Ser Ala Gln Asp Ser Leu
                                          10
      Gln Gln Thr Gly Gln Gln Met Lys Glu Lys Ala Gln Gly Ala Ala Asp
45
                                      25
      Val Val Lys Asp Lys Thr Gly Met Asn Lys Ser His
              35
      (2) INFORMATION FOR SEQ ID NO:84:
           (i) SEQUENCE CHARACTERISTICS:
50
                 (A) LENGTH: 43 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..43
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1018344
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:84:
60
      Met Asp Lys Ala Lys Asp Ala Ala Ala Ser Ala Gln Asp Ser Leu Gln
                                          10
      Gln Thr Gly Gln Gln Met Lys Glu Lys Ala Gln Gly Ala Ala Asp Val
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WO 00/40695 PCT/US00/00466 365 30 20 Val Lys Asp Lys Thr Gly Met Asn Lys Ser His 35 40 (2) INFORMATION FOR SEQ ID NO:85: 5 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 540 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 10 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..540 (D) OTHER INFORMATION: / Ceres Seq. ID 1018382 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:85: taaaagcaat aaaaaacaaa aatggcgttt actccgaaga tcatcacatg cctcattgtc 60 cttacqatct acatqacatc cccaacaqag tcaaccatcc agtgtgggac agtgacqagc 120 acactggcac agtgcgtgac ctacttgacc aacagtggtc cattgccatc acaatgctgc 180 qtgggagtca agtcattgta ccaattggct cagaccacac cggaccgtaa acaagtatgt 240 20 qaqtqcctta aactaqcqqq taaaqaaatc aaggqcctca acaccqacct tgtqqccqca 300 cttcctacca cttgtggtgt ttcaattccc taccccatca gttttagcac caattgcgac 360 agtatatcga ctgccqtqtg aaagaggcta gtgatcagat gtacgactaa tcaaacttgc 420 cagcttttaa cctaattaaa taaaagtatt ctgcttatat ttcccatttt atgattttat 480 cttcttatct atgtaaccac acgatttcat atgctaataa tgacaacgga tctttctctc 540 25 (2) INFORMATION FOR SEQ ID NO:86: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 119 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..119 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1018383 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:86: Met Ala Phe Thr Pro Lys Ile Ile Thr Cys Leu Ile Val Leu Thr Ile 10 Tyr Met Thr Ser Pro Thr Glu Ser Thr Ile Gln Cys Gly Thr Val Thr 40 20 25 Ser Thr Leu Ala Gln Cys Val Thr Tyr Leu Thr Asn Ser Gly Pro Leu 40 Pro Ser Gln Cys Cys Val Gly Val Lys Ser Leu Tyr Gln Leu Ala Gln 55 45 Thr Thr Pro Asp Arg Lys Gln Val Cys Glu Cys Leu Lys Leu Ala Gly 70 75 Lys Glu Ile Lys Gly Leu Asn Thr Asp Leu Val Ala Ala Leu Pro Thr 90 85 Thr Cys Gly Val Ser Ile Pro Tyr Pro Ile Ser Phe Ser Thr Asn Cys 50 105 100 Asp Ser Ile Ser Thr Ala Val 115 (2) INFORMATION FOR SEQ ID NO:87: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 102 amino acids

- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- 60 (ix) FEATURE:
  - (A) NAME/KEY: peptide
  - (B) LOCATION: 1..102

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(D) OTHER INFORMATION: / Ceres Seq. ID 1018384
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:87:
     Met Thr Ser Pro Thr Glu Ser Thr Ile Gln Cys Gly Thr Val Thr Ser
                                          10
 5
     Thr Leu Ala Gln Cys Val Thr Tyr Leu Thr Asn Ser Gly Pro Leu Pro
                  20
                                      25
      Ser Gln Cys Cys Val Gly Val Lys Ser Leu Tyr Gln Leu Ala Gln Thr
                                  40
      Thr Pro Asp Arg Lys Gln Val Cys Glu Cys Leu Lys Leu Ala Gly Lys
10
                              55
      Glu Ile Lys Gly Leu Asn Thr Asp Leu Val Ala Ala Leu Pro Thr Thr
                                              75
      Cys Gly Val Ser Ile Pro Tyr Pro Ile Ser Phe Ser Thr Asn Cys Asp
                      85
15
      Ser Ile Ser Thr Ala Val
                  100
      (2) INFORMATION FOR SEQ ID NO:88:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 75 amino acids
20
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
25
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 1018385
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:88:
      Lys Ala Ile Lys Asn Lys Asn Gly Val Tyr Ser Glu Asp His His Met
30
                                           10
      Pro His Cys Pro Tyr Asp Leu His Asp Ile Pro Asn Arg Val Asn His
                                      25
      Pro Val Trp Asp Ser Asp Glu His Thr Gly Thr Val Arg Asp Leu Leu
                                  40
35
      Asp Gln Gln Trp Ser Ile Ala Ile Thr Met Leu Arg Gly Ser Gln Val
                              55
      Ile Val Pro Ile Gly Ser Asp His Thr Gly Pro
                          70
      (2) INFORMATION FOR SEQ ID NO:89:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 844 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..844
                (D) OTHER INFORMATION: / Ceres Seq. ID 1020666
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:89:
      ctaaatattt aaagetttee tteaceagea ttgtattgee teeetggaae aeegaagete
                                                                              60
                                                                             120
      tggattaacc tattttcacc ttctaaggtt cttcgtcgtt cttattgatc tgtgaagatg
      tattcgggtc agaacaagat ccacaaggat aagggtgttg caccaacaga gtttgaagag
                                                                             180
                                                                             240
      caagttactc aggetttgtt tgacttggag aacaccaacc aggagttgaa aagcgagttg
55
                                                                             300
      aaagatctct acattaacca agctqttcaq atqqatattt ctqqcaaccg caaagctqtt
                                                                             360
      gtgatctacg ttccattcag gctgaggaaa gctttccgca agattcatct tcgtctggtc
      agagagettg agaagaagtt cagtggaaaa gatgtgatet ttgttgetae cagaagaate
                                                                             420
      atgegeceae caaaqaaagg ctcagetqtt caqagaceae geaacagaae tettaettet
                                                                             480
      gtccatgaag ccatgctcga ggatgtcgct taccctgctg agattgtagg aaagagaacc
                                                                             540
60
      agataccgtc ttgatggcac caagatcatg aaggtatttt tggatagtaa gctaaagaac
                                                                             600
      gacactgagt acaagctgga gacaatggtg ggtgtgtaca gaaaacttac agggaaagat
                                                                             660
      gtagttttcg agtacccagt catagaagct tgaagagatg aatgtttgtt tcattctttt
                                                                             720
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367

ttgtcggata gagagctttt gattctgttt gggagtaaaa ctctttgcaa agattttgtt 780 atttcagcat ttcgcaatct cttcttcaaa gttcgattta tcgatacaat tctttttcc 840 (2) INFORMATION FOR SEQ ID NO:90: 5 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 191 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 10 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..191 (D) OTHER INFORMATION: / Ceres Seq. ID 1020667 1.5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:90: Met Tyr Ser Gly Gln Asn Lys Ile His Lys Asp Lys Gly Val Ala Pro 5 10 Thr Glu Phe Glu Glu Gln Val Thr Gln Ala Leu Phe Asp Leu Glu Asn 25 20 Thr Asn Gln Glu Leu Lys Ser Glu Leu Lys Asp Leu Tyr Ile Asn Gln 40 Ala Val Gln Met Asp Ile Ser Gly Asn Arg Lys Ala Val Val Ile Tyr 55 Val Pro Phe Arg Leu Arg Lys Ala Phe Arg Lys Ile His Leu Arg Leu 25 70 75 Val Arg Glu Leu Glu Lys Lys Phe Ser Gly Lys Asp Val Ile Phe Val 90 8.5 Ala Thr Arg Arg Ile Met Arg Pro Pro Lys Lys Gly Ser Ala Val Gln 105 100 110 30 Arg Pro Arg Asn Arg Thr Leu Thr Ser Val His Glu Ala Met Leu Glu 120 125 115 Asp Val Ala Tyr Pro Ala Glu Ile Val Gly Lys Arg Thr Arg Tyr Arg 135 Leu Asp Gly Thr Lys Ile Met Lys Val Phe Leu Asp Ser Lys Leu Lys 35 155 150 Asn Asp Thr Glu Tyr Lys Leu Glu Thr Met Val Gly Val Tyr Arg Lys 170 165 Leu Thr Gly Lys Asp Val Val Phe Glu Tyr Pro Val Ile Glu Ala 180 185 40 (2) INFORMATION FOR SEQ ID NO:91: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 140 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 45 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..140 50 (D) OTHER INFORMATION: / Ceres Seq. ID 1020668 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:91: Met Asp Ile Ser Gly Asn Arg Lys Ala Val Val Ile Tyr Val Pro Phe 10 Arg Leu Arg Lys Ala Phe Arg Lys Ile His Leu Arg Leu Val Arg Glu 55 20 25 Leu Glu Lys Lys Phe Ser Gly Lys Asp Val Ile Phe Val Ala Thr Arg 40 Arg Ile Met Arg Pro Pro Lys Lys Gly Ser Ala Val Gln Arg Pro Arg 55 60 Asn Arg Thr Leu Thr Ser Val His Glu Ala Met Leu Glu Asp Val Ala 70 75 Tyr Pro Ala Glu Ile Val Gly Lys Arg Thr Arg Tyr Arg Leu Asp Gly

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368
                      85
                                          90
      Thr Lys Ile Met Lys Val Phe Leu Asp Ser Lys Leu Lys Asn Asp Thr
                                      105
      Glu Tyr Lys Leu Glu Thr Met Val Gly Val Tyr Arg Lys Leu Thr Gly
 5
              115
                                  120
      Lys Asp Val Val Phe Glu Tyr Pro Val Ile Glu Ala
          130
                              135
      (2) INFORMATION FOR SEQ ID NO:92:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 789 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
15
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..789
                (D) OTHER INFORMATION: / Ceres Seq. ID 1020784
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:92:
20
      agggttaggt ttgtcttcaa aatcgtcaga agaagaagaa gaagtcaaag acgataacaa
                                                                              60
      tgacqtcatc atcqtacctc cgatttqcqa tcqccqtcqt aqctttcctc tcaatcacca
                                                                             120
      ccatcaccac cgcacgtcct tgcaaaacct tcctaatctc ctcatattcc ctctcaataa
                                                                             180
      ccccagaaaa ccctaacctc gaatccgatt tcacctccac acgattcatc accgttttca
                                                                             240
      caatccgtcg tctaaatcct caccacgtgg ttcctttctt cgtcaatcgt cgtcacgaga
                                                                             300
25
      agccacagat ccaatcggat cgttccctcc ctttgatctc tgataacatc aattcgttta
                                                                             360
      gagategaae cagagatatt ettagegteg tegtegeget titgtteggt gitggttgtg
                                                                             420
      gtgctttaac cgctgctact atgtatcttg tttgggctct tgttgtgaat cgtcaaagct
                                                                             480
      atgattttga ggaagaagaa gatgattatg agaatgatga atctgatgct gctagcttga
                                                                             540
      agaaattggg ttatgttaag attccagctc cagctcctgc tcctgtgaag gaagctgctt
                                                                             600
30
      gatttgaggt tgtgaatttc agtttctgca tattatcatg atttacgttg ttgcttatct
                                                                             660
      atgttgttgc gtacaagtaa attcgtctct gttatggatt tatctaaact tactatactc
                                                                             720
      taagtatcag tgtttttacg atttgtagtt gatgtttctt atatatata ataagtgtga
                                                                             780
      ttatgatgc
      (2) INFORMATION FOR SEQ ID NO:93:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 180 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..180
                (D) OTHER INFORMATION: / Ceres Seq. ID 1020785
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:93:
      Met Thr Ser Ser Ser Tyr Leu Arg Phe Ala Ile Ala Val Val Ala Phe
                      5
                                           10
                                                               15
      Leu Ser Ile Thr Thr Ile Thr Thr Ala Arg Pro Cys Lys Thr Phe Leu
                                       25
50
      Ile Ser Ser Tyr Ser Leu Ser Ile Thr Pro Glu Asn Pro Asn Leu Glu
                                  40
                                                       45
      Ser Asp Phe Thr Ser Thr Arg Phe Ile Thr Val Phe Thr Ile Arg Arg
                              55
      Leu Asn Pro His His Val Val Pro Phe Phe Val Asn Arg Arg His Glu
55
                          70
                                               75
      Lys Pro Gln Ile Gln Ser Asp Arg Ser Leu Pro Leu Ile Ser Asp Asn
                                           90
      Ile Asn Ser Phe Arg Asp Arg Thr Arg Asp Ile Leu Ser Val Val
                  100
                                       105
                                                           110
60
      Ala Leu Leu Phe Gly Val Gly Cys Gly Ala Leu Thr Ala Ala Thr Met
                                   120
                                                       125
```

Tyr Leu Val Trp Ala Leu Val Val Asn Arg Gln Ser Tyr Asp Phe Glu

```
135
          130
     Glu Glu Glu Asp Asp Tyr Glu Asn Asp Glu Ser Asp Ala Ala Ser Leu
                                   155
                    150
     Lys Lys Leu Gly Tyr Val Lys Ile Pro Ala Pro Ala Pro Ala Pro Val
 5
                                          170
                      165
     Lys Glu Ala Ala
                 180
      (2) INFORMATION FOR SEQ ID NO:94:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 765 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
15
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1...765
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021525
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:94:
20
      acceteacte actittecqq titeaaccat taqcacaaaa aatetqaqet ecqaccaaaa
                                                                             60
      attggggaac gacgatggcg acgactette agaaactete eteteaaate cacegactet
                                                                            120
      ctccattcac cagateteta ategttegea cateegeaac gtetgeteet teteegtete
                                                                            180
      tgggatcgaa gaaagtctct gaccgaatcg tcaagctttc tgcgatcgat cccgatggat
      acaagcagga catcattgga ctctctggcc agactctcct tcgtgcgctc actcacactg
                                                                            300
25
      gtctgataga tccagcatcg catagattgg atgacatcga ggcttgctca gcggagtgcg
      aggttcagat cqcaqaqqaa tqqctcqaqa agctcccqcc qcqtacttac gatqaqqaqt
      atgtgctgaa gcgtagctct agatcccgta tcttgaacaa gcactcgcgt cttggttgcc
      aagttgtgtt aacgcaggag cttcaaggta tggtcgtcgc tgtccctgaa gctaagcctt
      gggatattcc gtaagttttt gttcttgagg atctcataaa tgttgggttc gttctcaatt
30
      tggctaataa aattcgcttt ggaaatttcg gtactctgaa gataactcca tttgaatctg
      taattttcgt ttgcttacat ttaaattttg taatgtggta aagaactctg cttgtttaaa
      cgattccctg cttcaagtat gggtgaatgt tgttttcttg attcc
      (2) INFORMATION FOR SEQ ID NO:95:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 159 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..159
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021526
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:95:
45
      Met Ala Thr Thr Leu Gln Lys Leu Ser Ser Gln Ile His Arg Leu Ser
                                          10
      Pro Phe Thr Arg Ser Leu Ile Val Arg Thr Ser Ala Thr Ser Ala Pro
                  20
                                      25
      Ser Pro Ser Leu Gly Ser Lys Lys Val Ser Asp Arg Ile Val Lys Leu
50
                                  40
      Ser Ala Ile Asp Pro Asp Gly Tyr Lys Gln Asp Ile Ile Gly Leu Ser
                              55
      Gly Gln Thr Leu Leu Arg Ala Leu Thr His Thr Gly Leu Ile Asp Pro
                                              75
55
      Ala Ser His Arg Leu Asp Asp Ile Glu Ala Cys Ser Ala Glu Cys Glu
                                           90
      Val Gln Ile Ala Glu Glu Trp Leu Glu Lys Leu Pro Pro Arg Thr Tyr
                                      105
                                                          110
                  100
      Asp Glu Glu Tyr Val Leu Lys Arg Ser Ser Arg Ser Arg Ile Leu Asn
60
                                                      125
                                  120
      Lys His Ser Arg Leu Gly Cys Gln Val Val Leu Thr Gln Glu Leu Gln
                              135
                                                  140
```

370 Gly Met Val Val Ala Val Pro Glu Ala Lys Pro Trp Asp Ile Pro 145 150 (2) INFORMATION FOR SEQ ID NO:96: (i) SEQUENCE CHARACTERISTICS: 5 (A) LENGTH: 588 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 10 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..588 (D) OTHER INFORMATION: / Ceres Seq. ID 1021563 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:96: 15 atcttcggaa agtctcattt ctcgatcccc aattcgtgga ttagggttaa aagaaccatt 60 tttattctcg tcgcgcaaca acaaatccag atcgaaaaag gaagaagaga tcgaaatggc 120 tttgagaagg gtttacagtg aaatcagagg gaagaaggtg acggagcttc caggctatat 180 caaatcgact ttttcaatgg agaccgtgaa gacctctgtg aagagaggac tcgataacta 240 caacgaaaaa tacattcaga ccagctccgt tgatcctatc cttcatatct gcttctacgg 300 20 catggctttc tcttaccttg tcgctctccc taatgagcgt cgccatcttg agcatcagca 360 gcatgctaag gagcacggtg gtcattgatc tcgtgggaat cgtttcgatc tcgagatgat 420 tttagggggg ttgctgtgaa atctttctct gcttgatggt gacgacgact caagaattgt 480 gtcttattgt ttcgttttct tgaattttcc tggataatgt tgacctaaag gaaaaccttt 540 ctttcgaatt acactccatg atagtcaata attgaagcat catgatgc 25 (2) INFORMATION FOR SEQ ID NO:97: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 128 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..128 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1021564 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:97: Ser Ser Glu Ser Leu Ile Ser Arg Ser Pro Ile Arg Gly Leu Gly Leu 5 10 Lys Glu Pro Phe Leu Phe Ser Ser Arg Asn Asn Lys Ser Arg Ser Lys 40 25 2.0 Lys Glu Glu Glu Ile Glu Met Ala Leu Arg Arg Val Tyr Ser Glu Ile 40 Arg Gly Lys Lys Val Thr Glu Leu Pro Gly Tyr Ile Lys Ser Thr Phe 55 45 Ser Met Glu Thr Val Lys Thr Ser Val Lys Arg Gly Leu Asp Asn Tyr 70 75 Asn Glu Lys Tyr Ile Gln Thr Ser Ser Val Asp Pro Ile Leu His Ile 90 Cys Phe Tyr Gly Met Ala Phe Ser Tyr Leu Val Ala Leu Pro Asn Glu 50 105 Arg Arg His Leu Glu His Gln Gln His Ala Lys Glu His Gly Gly His 120 115 (2) INFORMATION FOR SEQ ID NO:98: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 90 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide

60 (ix) FEATURE:

(A) NAME/KEY: peptide

(B) LOCATION: 1..90

WO 00/40695 PCT/US00/00466 371 (D) OTHER INFORMATION: / Ceres Seq. ID 1021565 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:98: Met Ala Leu Arg Arg Val Tyr Ser Glu Ile Arg Gly Lys Lys Val Thr 10 Glu Leu Pro Gly Tyr Ile Lys Ser Thr Phe Ser Met Glu Thr Val Lys 25 Thr Ser Val Lys Arg Gly Leu Asp Asn Tyr Asn Glu Lys Tyr Ile Gln 40 Thr Ser Ser Val Asp Pro Ile Leu His Ile Cys Phe Tyr Gly Met Ala 10 55 Phe Ser Tyr Leu Val Ala Leu Pro Asn Glu Arg Arg His Leu Glu His 70 75 Gln Gln His Ala Lys Glu His Gly Gly His 85 15 (2) INFORMATION FOR SEQ ID NO:99: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 586 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..586 25 (D) OTHER INFORMATION: / Ceres Seq. ID 1021576 (xi) SEQUENCE DESCRIPTION: SEO ID NO:99: ttgcctaaat cattttaaag actgtataga ggaaaacaaa actgcaaaaa caaaaataaa aaaaacatcg cacaagaaaa taaaagattt gtagaatcaa ctaagaaaat ggctagcact atgatgacta cattgcctca gttcaatggt cttcgagcca ccaaaatctc tgcagctcct 30 gtacaaggcc tggcaagtgt tcagcccatg agacgcaagg gaaatggagc tttgggtgca 240 aagtgtgact tcatcggttc atcaacaaat ctgataatgg taacgtcgac gaccctgatg 300 ttgttcgcgg ggagattcgg acttgcgcca tcagccaata ggaaggcgac agctggactt 360 aggttggagg cacgtgactc aggtctacaa acgggtgacc cggccgggtt cacgcttgcg 420 gacactttgg cttgtggcac cgttggtcat atcatcggtg taggagttgt ccttggcctt 480 35 aaaaacattg gtgctatttg aagttcctaa agctctttta tttgtatttg taaaatttgt 540 agatttttat aacaatattc tcatgcacct gaacgagatc taatgc (2) INFORMATION FOR SEQ ID NO:100: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 130 amino acids 40 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 45 (A) NAME/KEY: peptide (B) LOCATION: 1..130 (D) OTHER INFORMATION: / Ceres Seq. ID 1021577 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:100: Met Ala Ser Thr Met Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg 50 10 Ala Thr Lys Ile Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe 40 55 Ile Gly Ser Ser Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met Leu Phe Ala Gly Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala

75

90

Thr Ala Gly Leu Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly

Asp Pro Ala Gly Phe Thr Leu Ala Asp Thr Leu Ala Cys Gly Thr Val 105

85

100

372 Gly His Ile Ile Gly Val Gly Val Leu Gly Leu Lys Asn Ile Gly 115 120 Ala Ile 130 5 (2) INFORMATION FOR SEQ ID NO:101: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 126 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..126 15 (D) OTHER INFORMATION: / Ceres Seq. ID 1021578 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:101: Met Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg Ala Thr Lys Ile 10 Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg 20 20 25 Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe Ile Gly Ser Ser 40 Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met Leu Phe Ala Gly 55 25 Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala Thr Ala Gly Leu Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly Asp Pro Ala Gly 90 Phe Thr Leu Ala Asp Thr Leu Ala Cys Gly Thr Val Gly His Ile Ile 30 100 105 Gly Val Gly Val Val Leu Gly Leu Lys Asn Ile Gly Ala Ile 115 (2) INFORMATION FOR SEQ ID NO:102: (i) SEQUENCE CHARACTERISTICS: 35 (A) LENGTH: 125 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 40 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..125 (D) OTHER INFORMATION: / Ceres Seq. ID 1021579 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:102: 45 Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg Ala Thr Lys Ile Ser 10 Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe Ile Gly Ser Ser Thr 50 40 Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met Leu Phe Ala Gly Arg 55 Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala Thr Ala Gly Leu Arg 70 55 Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly Asp Pro Ala Gly Phe 85 90 Thr Leu Ala Asp Thr Leu Ala Cys Gly Thr Val Gly His Ile Ile Gly 100 105 Val Gly Val Val Leu Gly Leu Lys Asn Ile Gly Ala Ile 60 120 (2) INFORMATION FOR SEQ ID NO:103:

(i) SEQUENCE CHARACTERISTICS:

373

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(A) LENGTH: 801 base pairs(B) TYPE: nucleic acid
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- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- 5 (ii) MOLECULE TYPE: DNA (genomic)
  - (ix) FEATURE:
    - (A) NAME/KEY: -
    - (B) LOCATION: 1..801
  - (D) OTHER INFORMATION: / Ceres Seq. ID 1021927

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:103: atttttctct cagaatccat aaaaagagag agagataaat aaagagaaaa ctgaagaagc 60 tagaagatgg agaaaagtaa tgatcatgac aaggctagcc acggcggctc cggtggtggt 120 gccacggaga agtgggagga gactagcctc ggaatccgaa ctgccgagac aatgctccgg ttagctccgg tgggtctttg tgttgcagcg cttgttgtca tgcttaagga ctctgagact 15 300 aatgagttcg gttcaatttc ttactccaat ctcacagcct ttaggtactt ggtgcacgca aatggaatat gtgcaggcta ctctcttcta tcagcagcca ttgcagcgat gcctcgttct 360 tcttcgacaa tgcctcgtgt ttggaccttc ttttgtctcg accagcttct gacctacctg 420 gttcttgctg ctggagctgt atcagctgag gttctatact tggcttacaa tggagactca 480 gccattactt ggagcgatgc atgtagctcc tacggcggtt tctgtcatag agccactgct 20 totgttataa toacattott tgtggtttgt ttotacatcg ttotctotot aatotoctot tataagetet ttaetegett tgateeteet teeattgttg acteegeeaa gaatetegaa gtcgctgtct tcggaagtta gatcctccat ttgctcttca agtacatcta gtttgtcatg 720 780 tttcaaatgt tgtgtgtttt taactttggt caagagaaag aatgcttatg tatttcctct

25 (2) INFORMATION FOR SEQ ID NO:104:

tgtttcattg cttttcttcc t

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 204 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
- 30 (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:

50

- (A) NAME/KEY: peptide
- (B) LOCATION: 1..204
- 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1021928
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:104:
    Met Glu Lys Ser Asn Asp His Asp Lys Ala Ser His Gly Gly Ser Gly

1 5 10 15
Gly Gly Ala Thr Glu Lys Trp Glu Glu Thr Ser Leu Gly Ile Arg Thr

40 20 25 30

Ala Glu Thr Met Leu Arg Leu Ala Pro Val Gly Leu Cys Val Ala Ala 35 40 45

Leu Val Val Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile 50 55 60

45 Ser Tyr Ser Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly
65 70 75 80

Ile Cys Ala Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro

85 90 95
Arg Ser Ser Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp

100 105 110

Gln Leu Leu Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu

Val Leu Tyr Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp 130 135 140

Ala Cys Ser Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val 145 150 155 160

Ile Ile Thr Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile
165 170 175

Ser Ser Tyr Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp 180 185 190

Ser Ala Lys Asn Leu Glu Val Ala Val Phe Gly Ser 195 200

```
(2) INFORMATION FOR SEQ ID NO:105:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 169 amino acids
               (B) TYPE: amino acid
 5
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
10
               (B) LOCATION: 1..169
               (D) OTHER INFORMATION: / Ceres Seq. ID 1021929
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:105:
     Met Leu Arg Leu Ala Pro Val Gly Leu Cys Val Ala Ala Leu Val Val
                                        10
     Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile Ser Tyr Ser
15
                                    25
     Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly Ile Cys Ala
                                40
     Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro Arg Ser Ser
20
                            55
     Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp Gln Leu Leu
                        70
     Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu Val Leu Tyr
                                        90
25
     Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp Ala Cys Ser
                100 105
     Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val Ile Ile Thr
                 120 125
     Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile Ser Ser Tyr
30
                135 140
     Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp Ser Ala Lys
                       150
                                          155
     Asn Leu Glu Val Ala Val Phe Gly Ser
                    165
35
     (2) INFORMATION FOR SEQ ID NO:106:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 153 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
40
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..153
45
               (D) OTHER INFORMATION: / Ceres Seq. ID 1021930
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:106:
     Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile Ser Tyr Ser
                                       10
     Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly Ile Cys Ala
50
                                    2.5
     Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro Arg Ser Ser
                                40
     Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp Gln Leu Leu
                            55
55
     Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu Val Leu Tyr
                        70
                                            75
     Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp Ala Cys Ser
                                        90
     Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val Ile Ile Thr
60
                                   105
     Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile Ser Ser Tyr
                                120
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PCT/US00/00466 WO 00/40695 375 Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp Ser Ala Lys 130 135 Asn Leu Glu Val Ala Val Phe Gly Ser 145 150 5 (2) INFORMATION FOR SEQ ID NO:107: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 602 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..602 15 (D) OTHER INFORMATION: / Ceres Seq. ID 1021945 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:107: atcqaqcqat qqatttqqca qaactqtqqq cqattttcqq acccqqattc tccqqcqccq ttttcqqaac cqqqtqqtqq qqtcqacqcc qtcqtttqca qttccatcca aqttcccttc qtccattacc ttcccqqcat attcqcttct ctcqqaqctt tqatqttcaa ttqcqtcaqa 20 aaaqaaqaca ttqattactc tccttacgac gaaggcqagt qqaqattqaa qctqtqqctt 240 ttcatagcqt atgtcqtaqc atttgtttcc ttagctqctt ctgttqqctt qctqattcaa 300 gattcggttg tgaagactgg gccttcaact tggactggtg tggctggtgt ctttcaatgt 360 gtatttgtat tgataagtgg gctaatgtat tggacatcgc actcagagta gcgcacaggc 420 acatacttca totocqactc agattcatac tgtgacaact tacaagttcc totogtttct 480 25 ctactatgta aatteetgtg tettgtettg teacettgge attgtgtgeg agatteetet 540 gttgcatcag tgtgtgtttc tgttaaaact ttgtgtgaat gttattgaaa atactccaaa 600 (2) INFORMATION FOR SEQ ID NO:108: (i) SEQUENCE CHARACTERISTICS: 30 (A) LENGTH: 136 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 35 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..136 (D) OTHER INFORMATION: / Ceres Seq. ID 1021946 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:108: 40 Ile Glu Arg Trp Ile Trp Gln Asn Cys Gly Arg Phe Ser Asp Pro Asp 10 Ser Pro Ala Pro Phe Ser Glu Pro Gly Gly Gly Val Asp Ala Val Val 25 20 Cys Ser Ser Ile Gln Val Pro Phe Val His Tyr Leu Pro Gly Ile Phe 45 40 Ala Ser Leu Gly Ala Leu Met Phe Asn Cys Val Arg Lys Glu Asp Ile 55 Asp Tyr Ser Pro Tyr Asp Glu Gly Glu Trp Arg Leu Lys Leu Trp Leu 75 70 50 Phe Ile Ala Tyr Val Val Ala Phe Val Ser Leu Ala Ala Ser Val Gly 8.5 90 Leu Leu Ile Gln Asp Ser Val Val Lys Thr Gly Pro Ser Thr Trp Thr 105 100 Gly Val Ala Gly Val Phe Gln Cys Val Phe Val Leu Ile Ser Gly Leu 55 120 Met Tyr Trp Thr Ser His Ser Glu

135

(A) LENGTH: 82 amino acids

(B) TYPE: amino acid
(C) STRANDEDNESS:

(2) INFORMATION FOR SEQ ID NO:109: (i) SEQUENCE CHARACTERISTICS:

130

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376
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
 5
                (B) LOCATION: 1..82
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021947
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:109:
     Met Phe Asn Cys Val Arg Lys Glu Asp Ile Asp Tyr Ser Pro Tyr Asp
                                         10
10
     Glu Gly Glu Trp Arg Leu Lys Leu Trp Leu Phe Ile Ala Tyr Val Val
                  20
                                      25
     Ala Phe Val Ser Leu Ala Ala Ser Val Gly Leu Leu Ile Gln Asp Ser
                                  40
     Val Val Lys Thr Gly Pro Ser Thr Trp Thr Gly Val Ala Gly Val Phe
15
                             55
      Gln Cys Val Phe Val Leu Ile Ser Gly Leu Met Tyr Trp Thr Ser His
                       70
                                              75
     Ser Glu
20
      (2) INFORMATION FOR SEQ ID NO:110:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 693 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..693
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022170
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:110:
     gagtegetet tettteette egecattitt tietttetag gitgeagagt taaaggagaa
                                                                              60
      ggttttcaat tagggttttg tagagagaaa gatgagccga ag ttgggaa taccggtgaa
                                                                             120
      gcttcttcac gaggcctcag gtcatatcgt gacggtggag ctaaagagcg gcgagcttta
                                                                             180
35
     cagaggaagt atgattgagt gtgaggataa ctggaactgt cagctcgagg atattactta
                                                                             240
                                                                             300
      taccgccaag gatggtaagg tatcacagct tgagcatgtc ttcattcgag gcagtaaagt
      caggittatg gicataccag acatteteaa acatgeteea atgiteaage ggitagatge
                                                                             360
      tagaatcaag ggaaagagct catcactggg tgttggcaga ggtagagctg caatgcgagg
                                                                             420
                                                                             480
      gaaaccggct gccgggcccg ggcgtggaac tggaggaagg ggagcggtac cacctgtgag
40
      gagatgattg attttacgct caacatgtaa gctttgattc tgggatagtc tttctgtgat
                                                                             540
      atccaggtga aagccgcctt tcttggttct gtcattaatg ctttagagaa cactgtaaaa
                                                                             600
      ccaaacaaga gagagatata tccagatgca ctagtagttc tgtattttgg gttaccaaga
      tgtttgaatc attgacaagt tttcgttgtg atc
      (2) INFORMATION FOR SEQ ID NO:111:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 131 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..131
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022171
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:111:
      Met Ser Arg Ser Leu Gly Ile Pro Val Lys Leu Heu His Glu Ala Ser
                      5
                                          10
      Gly His Ile Val Thr Val Glu Leu Lys Ser Gly Glu Leu Tyr Arg Gly
                  20
                                      25
60
      Ser Met Ile Glu Cys Glu Asp Asn Trp Asn Cys Gln Leu Glu Asp Ile
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Thr Tyr Thr Ala Lys Asp Gly Lys Val Ser Gln Leu Glu His Val Phe

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377
                             55
     Ile Arg Gly Ser Lys Val Arg Phe Met Val Ile Pro Asp Ile Leu Lys
                        70
                                           75
     His Ala Pro Met Phe Lys Arg Leu Asp Ala Arg Ile Lys Gly Lys Ser
 5
                                        90
     Ser Ser Leu Gly Val Gly Arg Gly Arg Ala Ala Met Arg Gly Lys Pro
                         105 110
                100
     Ala Ala Gly Pro Gly Arg Gly Thr Gly Gly Arg Gly Ala Val Pro Pro
           115
                       120
10
     Val Arg Arg
         130
      (2) INFORMATION FOR SEQ ID NO:112:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 98 amino acids
15
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
20
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..98
               (D) OTHER INFORMATION: / Ceres Seq. ID 1022172
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:112:
     Met Ile Glu Cys Glu Asp Asn Trp Asn Cys Gln Leu Glu Asp Ile Thr
25
                                        10
      Tyr Thr Ala Lys Asp Gly Lys Val Ser Gln Leu Glu His Val Phe Ile
                                 25
                 2.0
     Arg Gly Ser Lys Val Arg Phe Met Val Ile Pro Asp Ile Leu Lys His
                                40
30
     Ala Pro Met Phe Lys Arg Leu Asp Ala Arg Ile Lys Gly Lys Ser Ser
                            5.5
      Ser Leu Gly Val Gly Arg Gly Arg Ala Ala Met Arg Gly Lys Pro Ala
                         70
                                            75
      Ala Gly Pro Gly Arg Gly Thr Gly Gly Arg Gly Ala Val Pro Pro Val
35
                     8.5
     Arg Arg
      (2) INFORMATION FOR SEQ ID NO:113:
          (i) SEQUENCE CHARACTERISTICS:
40
               (A) LENGTH: 65 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
45
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..65
               (D) OTHER INFORMATION: / Ceres Seq. ID 1022173
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:113:
50
      Glu Ser Leu Phe Phe Pro Ser Ala Ile Phe Phe Leu Gly Cys Arg
              5
                                     10
      Val Lys Gly Glu Gly Phe Gln Leu Gly Phe Cys Arg Glu Lys Asp Glu
                                     25
      Pro Lys Phe Gly Asn Thr Gly Glu Ala Ser Ser Arg Gly Leu Arg Ser
55
                                40
      Tyr Arg Asp Gly Gly Ala Lys Glu Arg Arg Ala Leu Gln Arg Lys Tyr
                             55
      Asp
      65
60
      (2) INFORMATION FOR SEQ ID NO:114:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 706 base pairs

YO 00/40093 FC1/0300/00400

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(B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
 5
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..706
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022554
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:114:
10
     agaccaaaaa agcggttggt gtaagatccc aaactcacag attcccaaat aatagtaata
                                                                              60
      ctcttcctct tctcaactct caccagtcac cagcagatca tcggagatgg gcggagacgg
      aaaagttttc accttqtccq aqqttictca qcacaqtaqc qccaaggatt gttggatcqt
                                                                             180
      catcgacggc aaggtttatg atgtgacaaa gttcttggat gatcatcctg gtggtgatga
                                                                             240
      ggttatcttg acttctacag ggaaagatgc gaccgatgat ttcgaggatg tgggacatag
                                                                             300
15
      ttcgactgcg aaagccatgc tagatgagta ctatgtgggt gatattgaca cagctactgt
                                                                             360
      gccggttaaa gctaagtttg tgcctcctac gtcgacgaaa gccgtggcta ctcaggataa
                                                                             420
      gageteggat tttgttatta ageteettea gtteettgtt eeaettetaa tettaggett
                                                                             480
      ggctttcggc attcggtact acactaagac caaggctcct tcttcttgaa agattgaatt
                                                                             540
      gtggcctaag tgaattgctt gtgtacgacg tggttaaact ctgtctctac ctctgtttgt
                                                                             600
20
      cttgtaagac atcattatat tctctattgc aatcaactga tagacctttg atttttaagc
                                                                             660
      catctacctt tgttgtctta tatcaatcaa atccatttac aatatc
      (2) INFORMATION FOR SEQ ID NO:115:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 140 amino acids
25
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
30
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..140
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022555
          (xi) SEQUENCE DESCRIPTION. SEQ ID NO:115:
      Met Gly Gly Asp Gly Lys Val Phe Thr Leu Ser Glu Val Ser Gln His
35
                                          10
      Ser Ser Ala Lys Asp Cys Trp Ile Val Ile Asp Gly Lys Val Tyr Asp
                  20
                                      25
      Val Thr Lys Phe Leu Asp Asp His Pro Gly Gly Asp Glu Val Ile Leu
                                  40
40
      Thr Ser Thr Gly Lys Asp Ala Thr Asp Asp Phe Glu Asp Val Gly His
                              55
      Ser Ser Thr Ala Lys Ala Met Leu Asp Glu Tyr Tyr Val Gly Asp Ile
                          70
                                              75
      Asp Thr Ala Thr Val Pro Val Lys Ala Lys Phe Val Pro Pro Thr Ser
45
                                          90
                      85
      Thr Lys Ala Val Ala Thr Gln Asp Lys Ser Ser Asp Phe Val Ile Lys
                                      105
                                                           110
                  100
      Leu Leu Gln Phe Leu Val Pro Leu Leu Ile Leu Gly Leu Ala Phe Gly
                                  120
50
      Ile Arg Tyr Tyr Thr Lys Thr Lys Ala Pro Ser Ser
                              135
      (2) INFORMATION FOR SEQ ID NO:116:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 70 amino acids
55
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
60
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..70
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022556
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:116:
     Met Leu Asp Glu Tyr Tyr Val Gly Asp Ile Asp Thr Ala Thr Val Pro
                                          10
      Val Lys Ala Lys Phe Val Pro Pro Thr Ser Thr Lys Ala Val Ala Thr
 5
                 20
                                      25
      Gln Asp Lys Ser Ser Asp Phe Val Ile Lys Leu Gln Phe Leu Val
                                 40
      Pro Leu Leu Ile Leu Gly Leu Ala Phe Gly Ile Arg Tyr Tyr Thr Lys
                              55
10
      Thr Lys Ala Pro Ser Ser
                          70
      (2) INFORMATION FOR SEQ ID NO:117:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 722 base pairs
15
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
20
                (A) NAME/KEY: -
                (B) LOCATION: 1..722
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022594
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:117:
      aacgtcatca ttctgacttt tctcaaagaa gagaaatcaa atcaccaagt catcggagcg
                                                                             60
25
      atcacagaga agaagaaaac catggcggaa gaagagaaaa gcgtgatgaa gcaagtgatg
                                                                             120
      gtggcgattg atgaaagcga atgtagtaaa cqcqctcttc aatggacgct cgtgtatctc
                                                                             180
                                                                             240
      aaagatagcc tcgccgattc cgacatcatc ctcttcactg ctcaacctca tcttgatctc
      agctgcgtct atgcttcctc ttatggcgcc gctccgattg agcttataaa ctcattgcaa
                                                                             300
      gagagtcata agaacgctgg attgaatcgt cttgatgaag gaaccaaaat ttgtgctgag
                                                                             360
30
      actggggtta ctccaagaaa ggtgttggaa tttggaaatc ctaaagaagc gatatgtgaa
                                                                          420
      gctgctgaga agcttggtgt tgatatgctt gtggttggaa gccatggtaa aggagcatta
                                                                             480
      caaaggactt toottggaag tigttagcaat tactgtgtta acaatgotaa gtgcccagtt
                                                                             540
      cttgtggtga gaacaaaagc ttgaagacct cttctcatga aagagttcga caaccctggc
                                                                             600
      ttgatgtgtg tgtgttgtaa acatattgat aataatgctc tgcttggtta ctattgtttg
                                                                             660
35
      aagaaacttt gtgagtaaac ataatttatt ttgtttgata atgaaagagc attgccgtgt
                                                                             720
      (2) INFORMATION FOR SEQ ID NO:118:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 187 amino acids
40
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
45
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..187
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022595
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:118:
      Asn Val Ile Ile Leu Thr Phe Leu Lys Glu Glu Lys Ser Asn His Gln
50
                                          10
      Val Ile Gly Ala Ile Thr Glu Lys Lys Lys Thr Met Ala Glu Glu Glu
                  20
                                      2.5
      Lys Ser Val Met Lys Gln Val Met Val Ala Ile Asp Glu Ser Glu Cys
                                  40
55
      Ser Lys Arg Ala Leu Gln Trp Thr Leu Val Tyr Leu Lys Asp Ser Leu
          50
                              55
      Ala Asp Ser Asp Ile Ile Leu Phe Thr Ala Gln Pro His Leu Asp Leu
                          70
                                              75
      Ser Cys Val Tyr Ala Ser Ser Tyr Gly Ala Ala Pro Ile Glu Leu Ile
60
                                           90
      Asn Ser Leu Gln Glu Ser His Lys Asn Ala Gly Leu Asn Arg Leu Asp
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380 Glu Gly Thr Lys Ile Cys Ala Glu Thr Gly Val Thr Pro Arg Lys Val 120 Leu Glu Phe Gly Asn Pro Lys Glu Ala Ile Cys Glu Ala Ala Glu Lys 135 5 Leu Gly Val Asp Met Leu Val Val Gly Ser His Gly Lys Gly Ala Leu 155 150 Gln Arg Thr Phe Leu Gly Ser Val Ser Asn Tyr Cys Val Asn Asn Ala 170 165 Lys Cys Pro Val Leu Val Val Arg Thr Lys Ala 10 180 185 (2) INFORMATION FOR SEQ ID NO:119: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 160 amino acids (B) TYPE: amino acid 15 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 20 (B) LOCATION: 1..160 (D) OTHER INFORMATION: / Ceres Seq. ID 1022596 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:119: Met Ala Glu Glu Lys Ser Val Met Lys Gln Val Met Val Ala Ile 10 5 25 Asp Glu Ser Glu Cys Ser Lys Arg Ala Leu Gln Trp Thr Leu Val Tyr 20 25 Leu Lys Asp Ser Leu Ala Asp Ser Asp Ile Ile Leu Phe Thr Ala Gln 40 Pro His Leu Asp Leu Ser Cys Val Tyr Ala Ser Ser Tyr Gly Ala Ala 30 55 Pro Ile Glu Leu Ile Asn Ser Leu Gln Glu Ser His Lys Asn Ala Gly 70 75 Leu Asn Arg Leu Asp Glu Gly Thr Lys Ile Cys Ala Glu Thr Gly Val 85 90 35 Thr Pro Arg Lys Val Leu Glu Phe Gly Asn Pro Lys Glu Ala Ile Cys 105 Glu Ala Ala Glu Lys Leu Gly Val Asp Met Leu Val Val Gly Ser His 120 Gly Lys Gly Ala Leu Gln Arg Thr Phe Leu Gly Ser Val Ser Asn Tyr 40 135 Cys Val Asn Asn Ala Lys Cys Pro Val Leu Val Val Arg Thr Lys Ala 150 (2) INFORMATION FOR SEQ ID NO:120: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 152 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 50 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..152 (D) OTHER INFORMATION: / Ceres Seq. ID 1022597 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:120: 55 Met Lys Gln Val Met Val Ala Ile Asp Glu Ser Glu Cys Ser Lys Arg 5 10 Ala Leu Gln Trp Thr Leu Val Tyr Leu Lys Asp Ser Leu Ala Asp Ser 25 Asp Ile Ile Leu Phe Thr Ala Gln Pro His Leu Asp Leu Ser Cys Val 60 40 Tyr Ala Ser Ser Tyr Gly Ala Ala Pro Ile Glu Leu Ile Asn Ser Leu

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Gln Glu Ser His Lys Asn Ala Gly Leu Asn Arg Leu Asp Glu Gly Thr
      Lys Ile Cys Ala Glu Thr Gly Val Thr Pro Arg Lys Val Leu Glu Phe
 5
      Gly Asn Pro Lys Glu Ala Ile Cys Glu Ala Ala Glu Lys Leu Gly Val
                 100
                                     105
      Asp Met Leu Val Val Gly Ser His Gly Lys Gly Ala Leu Gln Arg Thr
                                  120
      Phe Leu Gly Ser Val Ser Asn Tyr Cys Val Asn Asn Ala Lys Cys Pro
10
                             135
                                                  140
      Val Leu Val Val Arg Thr Lys Ala
                          150
      (2) INFORMATION FOR SEQ ID NO:121:
           (i) SEQUENCE CHARACTERISTICS:
15
                (A) LENGTH: 610 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
20
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..610
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022621
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:121:
25
      aattttgtgt ggtaattaac tggtgagaga gttcaacatc caatctttct ctctctctct
                                                                             60
      atcttttate teteacacte ttaaaatget tettetatet eegatetetg cetetettee
                                                                             120
      accgtcgttt caccggggta atttgatccg gcgatcaatt aagccgttag gtagggttgt
                                                                             180
      tgcaaaggcc aaggacaaca ccgacactgg tggatttctg gagacggcag ctatagccgg
                                                                             240
      tggtttagtg tcgacgccgg ttattggatg gtcactctac actttgaaga ccacgggatg
                                                                             300
30
      tggcttacct cccggaccgg ccggtttaat cggagcgttg gaaggtgtga gctacttggt
                                                                             360
      ggtggttggc atcgtgggct ggtctttgta cactaaaaca aaaactgggt caggtctgcc
                                                                             420
      aaatgggcca tttggtttgt tgggtgcggt cgagggttta tcgtatctgt cggttctagc
                                                                             480
      cattettgtg gtgtttggta ttcagttett ggataatggg teggtteeag gteeaettee
                                                                             540
      tagtgaccag tgttttggtt aaccacgtgt aatgtcgtat aaataaagtg tcaaattaat
                                                                             600
35
      aattgctttc
      (2) INFORMATION FOR SEQ ID NO:122:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 158 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..158
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022622
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:122:
      Met Leu Leu Ser Pro Ile Ser Ala Ser Leu Pro Pro Ser Phe His
                   5
                                          10
50
      Arg Gly Asn Leu Ile Arg Arg Ser Ile Lys Pro Leu Gly Arg Val Val
                                      25
      Ala Lys Ala Lys Asp Asn Thr Asp Thr Gly Gly Phe Leu Glu Thr Ala
                                  40
      Ala Ile Ala Gly Gly Leu Val Ser Thr Pro Val Ile Gly Trp Ser Leu
55
      Tyr Thr Leu Lys Thr Thr Gly Cys Gly Leu Pro Pro Gly Pro Ala Gly
                          70
                                               75
      Leu Ile Gly Ala Leu Glu Gly Val Ser Tyr Leu Val Val Val Gly Ile
                                           90
60
      Val Gly Trp Ser Leu Tyr Thr Lys Thr Lys Thr Gly Ser Gly Leu Pro
                                       105
                                                           110
      Asn Gly Pro Phe Gly Leu Leu Gly Ala Val Glu Gly Leu Ser Tyr Leu
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115
                                  120
                                                       125
     Ser Val Leu Ala Ile Leu Val Val Phe Gly Ile Gln Phe Leu Asp Asn
                              135
     Gly Ser Val Pro Gly Pro Leu Pro Ser Asp Gln Cys Phe Gly
 5
                          150
      (2) INFORMATION FOR SEQ ID NO:123:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 872 base pairs
                (B) TYPE: nucleic acid
10
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
15
                (B) LOCATION: 1..872
                (D) OTHER INFORMATION: / Ceres Seq. ID 1024375
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:123:
      atcgaacatt aaaacctgat tttactgaaa ctgtgctgcg aaggagaagc cgcaaaccat
                                                                              60
      tttttgtcaa tctagggttc ttcttcgtac ttgcagattt cgcctccgtt ttcactacga
                                                                             120
20
      tctgtgaaga tgttctccgc tcagaacaag atcaataagg acaagaatgc tgagccaaca
                                                                             180
      gaatgcyagg agcaagttgc tcaggctttg tttgatttgg agaacactaa ccaggagttg
                                                                             240
      aaaagcgagt tgaaagatct ctacatcaac caagctgttc acatggatat ctctggaaac
                                                                             300
                                                                             360
      cgcaaagctg ttgtgattta cgttccattc agattgagga aagctttccg caagattcat
      ccccgtctcg tcagagagct tgagaagaag ttcagtggaa aggatgttat ctttgttacc
                                                                             420
25
      acaagaagga tcatgcgtcc ccccaagaag ggtgctgctg ttcagaggcc acgtaacaga
                                                                             480
                                                                             540
      actettacet cagtteatga agetatgett gaagatgttg ettteeeege tgagattgtt
      ggaaagcgta ctcgctaccg tcttgatggt tccaagatca tgaaggtctt tttggatgcc
                                                                             600
      aaggaaaaga acaacacaga gtacaagctc gagactatgg tcggtgtgta ccgtaaactt
                                                                             660
      actggcaaag atgttgtttt tgagtaccca gtcgaagctt gaaagaagat gatgaagaac
                                                                             720
30
                                                                             780
      catcaggata gtgaaagaga gcttttgttt atgttttgtg gtatttagga tgaaggaaac
                                                                             840
      tctcttgatt cagttccttg ttcacaatct ttaatgttct atttacaatg actacttttt
      gtgtttttca attttgaaac ctctatgatt tc
      (2) INFORMATION FOR SEQ ID NO:124:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 190 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..190
                (D) OTHER INFORMATION: / Ceres Seq. ID 1024376
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:124:
45
      Met Phe Ser Ala Gln Asn Lys Ile Asn Lys Asp Lys Asn Ala Glu Pro
                                           10
      Thr Glu Cys Glu Glu Gln Val Ala Gln Ala Leu Phe Asp Leu Glu Asn
                  2.0
                                       25
      Thr Asn Gln Glu Leu Lys Ser Glu Leu Lys Asp Leu Tyr Ile Asn Gln
50
      Ala Val His Met Asp Ile Ser Gly Asn Arg Lys Ala Val Val Ile Tyr
                               55
      Val Pro Phe Arg Leu Arg Lys Ala Phe Arg Lys Ile His Pro Arg Leu
                          70
                                               75
55
      Val Arg Glu Leu Glu Lys Lys Phe Ser Gly Lys Asp Val Ile Phe Val
                                           90
      Thr Thr Arg Arg Ile Met Arg Pro Pro Lys Lys Gly Ala Ala Val Gln
                                       105
                                                           110
      Arg Pro Arg Asn Arg Thr Leu Thr Ser Val His Glu Ala Met Leu Glu
60
                                   120
      Asp Val Ala Phe Pro Ala Glu Ile Val Gly Lys Arg Thr Arg Tyr Arg
          130
                               135
                                                   140
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383 Leu Asp Gly Ser Lys Ile Met Lys Val Phe Leu Asp Ala Lys Glu Lys 150 155 Asn Asn Thr Glu Tyr Lys Leu Glu Thr Met Val Gly Val Tyr Arg Lys 165 170 5 Leu Thr Gly Lys Asp Val Val Phe Glu Tyr Pro Val Glu Ala 180 185 (2) INFORMATION FOR SEQ ID NO:125: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 139 amino acids 10 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 15 (A) NAME/KEY: peptide (B) LOCATION: 1..139 (D) OTHER INFORMATION: / Ceres Seq. ID 1024377 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:125: Met Asp Ile Ser Gly Asn Arg Lys Ala Val Val Ile Tyr Val Pro Phe 20 10 5 Arg Leu Arg Lys Ala Phe Arg Lys Ile His Pro Arg Leu Val Arg Glu 20 25 30 Leu Glu Lys Lys Phe Ser Gly Lys Asp Val Ile Phe Val Thr Thr Arg 40 4.5 25 Arg Ile Met Arg Pro Pro Lys Lys Gly Ala Ala Val Gln Arg Pro Arg 55 60 Asn Arg Thr Leu Thr Ser Val His Glu Ala Met Leu Glu Asp Val Ala 70 75 Phe Pro Ala Glu Ile Val Gly Lys Arg Thr Arg Tyr Arg Leu Asp Gly 30 90 Ser Lys Ile Met Lys Val Phe Leu Asp Ala Lys Glu Lys Asn Asn Thr 105 Glu Tyr Lys Leu Glu Thr Met Val Gly Val Tyr Arg Lys Leu Thr Gly 115 120 35 Lys Asp Val Val Phe Glu Tyr Pro Val Glu Ala 130 135 (2) INFORMATION FOR SEQ ID NO:126: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 545 base pairs 40 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 45 (A) NAME/KEY: -(B) LOCATION: 1..545 (D) OTHER INFORMATION: / Ceres Seq. ID 1024535 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:126: aaaaattqtt aaqttcqtqt ccqcqaqatc aqaqtttqtq tccqqttata cqattcqtcq 60 50 gtaccaaaac cctccgagaa gtgtttcagg tgggaaggta tctctgtcta cacgatgtcc 120 ggcgaggaag aagccaccgt gagggagcca ctagatctga ttaggctgag tctcgacgag 180 agaatctatq tcaaqctccg gtcagaccgc gaacttcgcg gcaaqcttca cgcgtttgat 240 cagcatttga atatgattct gggtgatgtt gaagaaacta tcactacagt agaaatcgat 300 gacgagacat atgaagagat tgttcggact acaaagcgga cgattgagtt tctattcgtg 360 55 agaggagatg gagtgatatt ggtgtctcca ccgctgagga cagcagcctg agttcaaact 420 caaatctttt gtgtcctctt ttcgtaaggg gaatttgtga ctattcttgt atttagtcac 480 540 ttcttgtctt aactaaaaca agcttgaaga agctagtttg cttaaatttc caatgctttt cagac (2) INFORMATION FOR SEQ ID NO:127: 60 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 98 amino acids

(B) TYPE: amino acid

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(C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
5
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..98
               (D) OTHER INFORMATION: / Ceres Seq. ID 1024536
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:127:
     Met Ser Gly Glu Glu Glu Ala Thr Val Arg Glu Pro Leu Asp Leu Ile
10
                                        10
     Arg Leu Ser Leu Asp Glu Arg Ile Tyr Val Lys Leu Arg Ser Asp Arg
                                     25
     Glu Leu Arg Gly Lys Leu His Ala Phe Asp Gln His Leu Asn Met Ile
                                 4.0
15
     Leu Gly Asp Val Glu Glu Thr Ile Thr Thr Val Glu Ile Asp Asp Glu
                            55
     Thr Tyr Glu Glu Ile Val Arg Thr Thr Lys Arg Thr Ile Glu Phe Leu
                        70
                                     75
     Phe Val Arg Gly Asp Gly Val Ile Leu Val Ser Pro Pro Leu Arg Thr
20
                                         90
                     85
     Ala Ala
      (2) INFORMATION FOR SEQ ID NO:128:
          (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 53 amino acids
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..53
               (D) OTHER INFORMATION: / Ceres Seq. ID 1024537
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:128:
35
      Lys Leu Leu Ser Ser Cys Pro Arg Asp Gln Ser Leu Cys Pro Val Ile
                                        10
      Arg Phe Val Gly Thr Lys Thr Leu Arg Glu Val Phe Gln Val Gly Arg
                      25
      Tyr Leu Cys Leu His Asp Val Arg Arg Gly Arg Ser His Arg Glu Gly
40
             35
                                40
      Ala Thr Arg Ser Asp
          50
      (2) INFORMATION FOR SEQ ID NO:129:
           (i) SEQUENCE CHARACTERISTICS:
45
                (A) LENGTH: 52 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
50
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..52
                (D) OTHER INFORMATION: / Ceres Seq. ID 1024538
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:129:
55
      Met Ile Leu Gly Asp Val Glu Glu Thr Ile Thr Thr Val Glu Ile Asp
                                         10
      Asp Glu Thr Tyr Glu Glu Ile Val Arg Thr Thr Lys Arg Thr Ile Glu
                         25
      Phe Leu Phe Val Arg Gly Asp Gly Val Ile Leu Val Ser Pro Pro Leu
          35
60
                          40
      Arg Thr Ala Ala
          50
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(2) INFORMATION FOR SEQ ID NO:130:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 797 base pairs
                (B) TYPE: nucleic acid
5
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
10
                (B) LOCATION: 1..797
                (D) OTHER INFORMATION: / Ceres Seq. ID 1025683
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:130:
      aaccettaaa eteegttaee etaaaeetea ageegtegte teeaaagtte eeeeteagag
                                                                              60
      acqaaaaaat qaaqaccatt ttqtcctcqq aqactatqqa catccccqac qqcqtcqcca
                                                                             120
15
      ttaaggtaaa cgccaaggtg attgaggtcg aaggtccacg aggtaaactc actcgtgact
                                                                             180
      tcaaqcatct qaatctcqat ttccaqttqa ttaaaqacca aqtcactqqa aaacqtcaqc
                                                                             240
      ttaagattga ttcttggttt ggttctcgta agacaagtgc ttcgattaga actgctttaa
                                                                             300
      qccatqttqa taatctcatt qctqqtqtta ctcaaqqttt tctttataqa atqaqatttq
                                                                             360
      tqtatqctca ttttcctatc aatqcttcta ttqatqqtaa caataaqaqt attqaqattc
                                                                             420
20
      gtaacttect tggtgagaag aaggtgagga aggttgagat gttggatggt gttaagattg
                                                                             480
      ttcqatctqa qaaqqttaaq qatqaqatta ttcttqaqqq aaatqatatt qaqcttqttt
                                                                             540
      cacggtcttg tgctttgatc aatcagaaat gtcatgtgaa gaagaaggat atcaggaagt
                                                                             600
      ttcttgatgg tatctatgtg agcgagaaag gcaagatcgc agtcgaggaa tgaagttcgc
                                                                             660
      tattgaaagt totatattag attttatatg gtgaagcagt gatattatcg tagotttttg
                                                                             720
25
      ttttacatct ttcttatgga tttgtagttt tgtttttcct tttgaatcat tgatgaatge
                                                                             780
      aaattcatta tccgttc
      (2) INFORMATION FOR SEQ ID NO:131:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 216 amino acids
30
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..216
                (D) OTHER INFORMATION: / Ceres Seq. ID 1025684
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:131:
      Pro Leu Asn Ser Val Thr Leu Asn Leu Lys Pro Ser Ser Pro Lys Phe
40
                                           10
                                                               15
      Pro Leu Arg Asp Glu Lys Met Lys Thr Ile Leu Ser Ser Glu Thr Met
                                       25
      Asp Ile Pro Asp Gly Val Ala Ile Lys Val Asn Ala Lys Val Ile Glu
                                  40
                                                       4.5
45
      Val Glu Gly Pro Arg Gly Lys Leu Thr Arg Asp Phe Lys His Leu Asn
                              55
      Leu Asp Phe Gln Leu Ile Lys Asp Gln Val Thr Gly Lys Arg Gln Leu
                          70
                                               75
      Lys Ile Asp Ser Trp Phe Gly Ser Arg Lys Thr Ser Ala Ser Ile Arg
50
                      85
                                           90
      Thr Ala Leu Ser His Val Asp Asn Leu Ile Ala Gly Val Thr Gln Gly
                  100
                                       105
                                                           110
      Phe Leu Tyr Arg Met Arg Phe Val Tyr Ala His Phe Pro Ile Asn Ala
              115
                                  120
                                                       125
55
      Ser Ile Asp Gly Asn Asn Lys Ser Ile Glu Ile Arg Asn Phe Leu Gly
                              135
                                                   140
      Glu Lys Lys Val Arg Lys Val Glu Met Leu Asp Gly Val Lys Ile Val
                          150
                                               155
      Arg Ser Glu Lys Val Lys Asp Glu Ile Ile Leu Glu Gly Asn Asp Ile
60
                                          170
                                                              175
      Glu Leu Val Ser Arg Ser Cys Ala Leu Ile Asn Gln Lys Cys His Val
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185

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Lys Lys Lys Asp Ile Arg Lys Phe Leu Asp Gly Ile Tyr Val Ser Glu
         195 200 205
     Lys Gly Lys Ile Ala Val Glu Glu
         210
5
     (2) INFORMATION FOR SEQ ID NO:132:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 194 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
10
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..194
15
              (D) OTHER INFORMATION: / Ceres Seq. ID 1025685
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:132:
     Met Lys Thr Ile Leu Ser Ser Glu Thr Met Asp Ile Pro Asp Gly Val
                  5
                                     10
     Ala Ile Lys Val Asn Ala Lys Val Ile Glu Val Glu Gly Pro Arg Gly
20
         20
                                 25
     Lys Leu Thr Arg Asp Phe Lys His Leu Asn Leu Asp Phe Gln Leu Ile
                              40
     Lys Asp Gln Val Thr Gly Lys Arg Gln Leu Lys Ile Asp Ser Trp Phe
                           55
25
     Gly Ser Arg Lys Thr Ser Ala Ser Ile Arg Thr Ala Leu Ser His Val
     Asp Asn Leu Ile Ala Gly Val Thr Gln Gly Phe Leu Tyr Arg Met Arg
                                      90
     Phe Val Tyr Ala His Phe Pro Ile Asn Ala Ser Ile Asp Gly Asn Asn
30
                                  105
     Lys Ser Ile Glu Ile Arg Asn Phe Leu Gly Glu Lys Lys Val Arg Lys
                               120
     Val Glu Met Leu Asp Gly Val Lys Ile Val Arg Ser Glu Lys Val Lys
                           135
                                              140
35
     Asp Glu Ile Ile Leu Glu Gly Asn Asp Ile Glu Leu Val Ser Arg Ser
                       150
                                          155
     Cys Ala Leu Ile Asn Gln Lys Cys His Val Lys Lys Lys Asp Ile Arg
                   165 170 175
     Lys Phe Leu Asp Gly Ile Tyr Val Ser Glu Lys Gly Lys Ile Ala Val
40
                180
                                  185
     Glu Glu
      (2) INFORMATION FOR SEQ ID NO:133:
          (i) SEQUENCE CHARACTERISTICS:
45
               (A) LENGTH: 185 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
50
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..185
               (D) OTHER INFORMATION: / Ceres Seq. ID 1025686
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:133:
55
     Met Asp Ile Pro Asp Gly Val Ala Ile Lys Val Asn Ala Lys Val Ile
                                      10
      Glu Val Glu Gly Pro Arg Gly Lys Leu Thr Arg Asp Phe Lys His Leu
              20
                                  25
      Asn Leu Asp Phe Gln Leu Ile Lys Asp Gln Val Thr Gly Lys Arg Gln
60
       35 40
      Leu Lys Ile Asp Ser Trp Phe Gly Ser Arg Lys Thr Ser Ala Ser Ile
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387
     Arg Thr Ala Leu Ser His Val Asp Asn Leu Ile Ala Gly Val Thr Gln
                          70
                                              75
     Gly Phe Leu Tyr Arg Met Arg Phe Val Tyr Ala His Phe Pro Ile Asn
                                          90
 5
     Ala Ser Ile Asp Gly Asn Asn Lys Ser Ile Glu Ile Arg Asn Phe Leu
                                      105
      Gly Glu Lys Lys Val Arg Lys Val Glu Met Leu Asp Gly Val Lys Ile
                                  120
                                                      125
      Val Arg Ser Glu Lys Val Lys Asp Glu Ile Ile Leu Glu Gly Asn Asp
10
                              135
                                                  140
      Ile Glu Leu Val Ser Arg Ser Cys Ala Leu Ile Asn Gln Lys Cys His
                         150
                                              155
      Val Lys Lys Lys Asp Ile Arg Lys Phe Leu Asp Gly Ile Tyr Val Ser
                                          170
                     165
15
      Glu Lys Gly Lys Ile Ala Val Glu Glu
                  180
      (2) INFORMATION FOR SEQ ID NO:134:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 555 base pairs
20
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
25
                (A) NAME/KEY: -
                (B) LOCATION: 1..555
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027152
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:134:
      ggttttgtca tcgtgctcct cttctcctcc accgcacaga tccaagacat tttgattaca
                                                                              60
30
      ctccatcgcc ggcgaaaatg gtgacggaaa agaaaacgaa gaagtctcat gagggtatca
                                                                             120
      acagtaggtt ggctcttgtt atgaagagtg gtaaatacac tcttggttac aagtctgttc
                                                                             180
      tcaaatccct ccgtggctcc aaagggaaac tgattctaat ctccaccaat tgcccaccgt
                                                                             240
      tgagaagatc agagattgag tactacgcaa tgctcgctaa agttggtgtc caccattaca
                                                                             300
                                                                             360
      atggaaataa cgttgatttg ggaacagctt gcggaaagta cttccgtgtt tcttgtctga
35
      gcattgttga tcctggtgat tctgacatta tcaagtcaat tcctggtgac cagtgataca
                                                                             420
      attttcagat tttattttc ccagactttt tacccgtctt tgtattcgag tctctcaaaa
                                                                             480
      ttttatgagc aatagacatt gtgtttcttc tttgtcaagt tttcttatat tgaggaccgg
                                                                             540
      aaaagaaatt tcttt
      (2) INFORMATION FOR SEQ ID NO:135:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 137 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..137
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1027153
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:135:
      Phe Cys His Arg Ala Pro Leu Leu His Arg Thr Asp Pro Arg His
                      5
                                          10
      Phe Asp Tyr Thr Pro Ser Pro Ala Lys Met Val Thr Glu Lys Lys Thr
                  20
                                      25
55
      Lys Lys Ser His Glu Gly Ile Asn Ser Arg Leu Ala Leu Val Met Lys
                                   4.0
      Ser Gly Lys Tyr Thr Leu Gly Tyr Lys Ser Val Leu Lys Ser Leu Arg
                              55
      Gly Ser Lys Gly Lys Leu Ile Leu Ile Ser Thr Asn Cys Pro Pro Leu
60
                          70
                                              75
      Arg Arg Ser Glu Ile Glu Tyr Tyr Ala Met Leu Ala Lys Val Gly Val
                                           90
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His His Tyr Asn Gly Asn Asn Val Asp Leu Gly Thr Ala Cys Gly Lys
     100 105 110

Tyr Phe Arg Val Ser Cys Leu Ser Ile Val Asp Pro Gly Asp Ser Asp
                   120
5
     Ile Ile Lys Ser Ile Pro Gly Asp Gln
                      135
         130
     (2) INFORMATION FOR SEQ ID NO:136:
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 112 amino acids
10
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
15
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..112
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027154
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:136:
     Met Val Thr Glu Lys Lys Thr Lys Lys Ser His Glu Gly Ile Asn Ser
20
                                          10
     Arg Leu Ala Leu Val Met Lys Ser Gly Lys Tyr Thr Leu Gly Tyr Lys
                                      25
     Ser Val Leu Lys Ser Leu Arg Gly Ser Lys Gly Lys Leu Ile Leu Ile
                                  40
25
      Ser Thr Asn Cys Pro Pro Leu Arg Arg Ser Glu Ile Glu Tyr Tyr Ala
                              55
     Met Leu Ala Lys Val Gly Val His His Tyr Asn Gly Asn Asn Val Asp
                          70
      Leu Gly Thr Ala Cys Gly Lys Tyr Phe Arg Val Ser Cys Leu Ser Ile
30
                      85
                                          90
      Val Asp Pro Gly Asp Ser Asp Ile Ile Lys Ser Ile Pro Gly Asp Gln
                  100
                                     105
      (2) INFORMATION FOR SEQ ID NO:137:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 91 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..91
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027155
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:137:
      Met Lys Ser Gly Lys Tyr Thr Leu Gly Tyr Lys Ser Val Leu Lys Ser
45
                                          10
      Leu Arg Gly Ser Lys Gly Lys Leu Ile Leu Ile Ser Thr Asn Cys Pro
                                      25
      Pro Leu Arg Arg Ser Glu Ile Glu Tyr Tyr Ala Met Leu Ala Lys Val
50
                                  40
      Gly Val His His Tyr Asn Gly Asn Asn Val Asp Leu Gly Thr Ala Cys
                             55
                                                  60
      Gly Lys Tyr Phe Arg Val Ser Cys Leu Ser Ile Val Asp Pro Gly Asp
                      70
55
      Ser Asp Ile Ile Lys Ser Ile Pro Gly Asp Gln
                     85
      (2) INFORMATION FOR SEQ ID NO:138:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 653 base pairs
60
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
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389
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..653
5
                (D) OTHER INFORMATION: / Ceres Seq. ID 1028095
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:138:
      aaataaagtt tttatttatc tatctatcca attggagacg aatcatatcg agcgatggat
                                                                              60
      ttggcagaac tgtgggcgat tttcggaccc ggattctccg gcgccgtttt cggaaccggg
                                                                             120
      tggtggtttt gggtcgacgc cgtcgtttgc agttccatcc aagttccctt cgtccattac
                                                                             180
10
      cttcccggca tattcqcttc tctcggagct ttgatgttca attgcgtcag aaaagaagac
                                                                             240
      attgattact ctccttacga cgaaggcgag tggagattga agctgtggct tttcatagcg
                                                                             300
      tatqtcqtaq catttqtttc cttaqctqct tctqttggct tqctqattca aqattcqqtt
                                                                             360
      qtqaaqactq qqccttcaac ttqqactqqt qtqqctggtq tctttcaatq tqtatttqta
                                                                             420
      ttgataagtg ggctaatgta ttggacatcg cactcagagt agcgcacagg cacatacttc
                                                                             480
15
      atctccgact cagattcata ctgtgacaac ttacaagttc ctctcgtttc tctactatgt
                                                                             540
      aaattcctqt qtcttqtctt qtcaccttqq cattqtqtqc qaqattcctc tqttqcatca
                                                                             600
      qtqtqttt ctqttaaaac tttqtqtqaa tqttattqaa aatactccaa atc
      (2) INFORMATION FOR SEQ ID NO:139:
           (i) SEQUENCE CHARACTERISTICS:
20
                (A) LENGTH: 135 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
25
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..135
                (D) OTHER INFORMATION: / Ceres Seq. ID 1028096
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:139:
30
      Met Asp Leu Ala Glu Leu Trp Ala Ile Phe Gly Pro Gly Phe Ser Gly
                    5
                                           10
      Ala Val Phe Gly Thr Gly Trp Trp Phe Trp Val Asp Ala Val Cys
                                      25
      Ser Ser Ile Gln Val Pro Phe Val His Tyr Leu Pro Gly Ile Phe Ala
35
      Ser Leu Gly Ala Leu Met Phe Asn Cys Val Arg Lys Glu Asp Ile Asp
      Tyr Ser Pro Tyr Asp Glu Gly Glu Trp Arg Leu Lys Leu Trp Leu Phe
                          70
40
      Ile Ala Tyr Val Val Ala Phe Val Ser Leu Ala Ala Ser Val Gly Leu
      Leu Ile Gln Asp Ser Val Val Lys Thr Gly Pro Ser Thr Trp Thr Gly
                                      105
                                                           110
                  1.00
      Val Ala Gly Val Phe Gln Cys Val Phe Val Leu Ile Ser Gly Leu Met
45
                                  120
              115
      Tyr Trp Thr Ser His Ser Glu
                              135
          130
      (2) INFORMATION FOR SEQ ID NO:140:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 82 amino acids
                 (B) TYPE: amino acid
                (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..82
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1028097
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:140:
60
      Met Phe Asn Cys Val Arg Lys Glu Asp Ile Asp Tyr Ser Pro Tyr Asp
                                          10
      Glu Gly Glu Trp Arg Leu Lys Leu Trp Leu Phe Ile Ala Tyr Val Val
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- 10 (2) INFORMATION FOR SEQ ID NO:141:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 80 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:

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- (A) NAME/KEY: peptide
- (B) LOCATION: 1..80
- 20 (D) OTHER INFORMATION: / Ceres Seq. ID 1028098
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:141:

Ile Lys Phe Leu Phe Ile Tyr Leu Ser Asn Trp Arg Arg Ile Ile Ser 1 5 10 15

Arg Arg Phe Arg Asn Arg Val Val Leu Gly Arg Arg Arg 35 40 45

Leu Gln Phe His Pro Ser Ser Leu Arg Pro Leu Pro Ser Arg His Ile
50 55 60

- Arg Phe Ser Arg Ser Phe Asp Val Gln Leu Arg Gln Lys Arg Arg His 65 70 75 80
  - (2) INFORMATION FOR SEQ ID NO:142:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 779 base pairs
      - (B) TYPE: nucleic acid
      - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
      (ii) MOLECULE TYPE: DNA (genomic)
    - (ix) FEATURE:
- 40 (A) NAME/KEY: -
  - (B) LOCATION: 1..779
  - (D) OTHER INFORMATION: / Ceres Seq. ID 1028608
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:142:

gtctaaactc taaactctaa agactttgtc atttcgtgaa ccgtcgtcat aactgatcct 60 45 tocaaagaat otocaaaato ootaaactto gaaaatggca gaaaccgcog agacaatcaa 120 caccacgate teategoege caceggaate egaaagetee accaetattt etgeaatgae 180 agatecaaca teteaagaag eageeteaaa agaeacagat etgaegaaag aageegaate 240 agagaagaaa ccaggaggaa teteteteeg aatetggeea eegaeteaga aaaetegega 300 cgccgttctg aatcgcctga tcgagacctt atccaccgaa tcaatcctct ctaagagata 360 50 cggtactctt aaatctgacg atgcaaccac cgtcgcgaaa cttatcgaag aagaggctta 420 tggtgttgct tcgaatgctg tgtcgagtga tgatgatggg attaagattc ttgagcttta 480 ttctaaagag attagtaaga ggatgcttga atctgttaag gctagatcta atgctagtgt 540 tggaaatgga agtgtggagg atgctaatac tgatgctagt gaggtttcta aagatgatgc 600 tggtcctggt tcagaggagg agaagagtga ggcttgagaa atttggtttt acttccttat 660 55 tataatctgt tgtctcttag ggttttgctg ttcttgtaat gatgatgatg acttgtggat 720 ttggtattat gcatatttga agatgtttct tcttccttaa tatgagtgca agtgtcttc

- (2) INFORMATION FOR SEQ ID NO:143:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 180 amino acids
  - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear

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(ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..180
 5
               (D) OTHER INFORMATION: / Ceres Seq. ID 1028609
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:143:
     Met Ala Glu Thr Ala Glu Thr Ile Asn Thr Thr Ile Ser Ser Pro Pro
                                    10
     Pro Glu Ser Glu Ser Ser Thr Thr Ile Ser Ala Met Thr Asp Pro Thr
10
                                    25
     Ser Gln Glu Ala Ala Ser Lys Asp Thr Asp Leu Thr Lys Glu Ala Glu
                                40
     Ser Glu Lys Lys Pro Gly Gly Ile Ser Leu Arg Ile Trp Pro Pro Thr
                            55
15
     Gln Lys Thr Arg Asp Ala Val Leu Asn Arg Leu Ile Glu Thr Leu Ser
                     70
                                            75
     Thr Glu Ser Ile Leu Ser Lys Arg Tyr Gly Thr Leu Lys Ser Asp Asp
                    85
                                       90
     Ala Thr Thr Val Ala Lys Leu Ile Glu Glu Glu Ala Tyr Gly Val Ala
20
                100
                                   105
     Ser Asn Ala Val Ser Ser Asp Asp Gly Ile Lys Ile Leu Glu Leu
                                120
     Tyr Ser Lys Glu Ile Ser Lys Arg Met Leu Glu Ser Val Lys Ala Arg
                            135
                                               140
25
     Ser Asn Ala Ser Val Gly Asn Gly Ser Val Glu Asp Ala Asn Thr Asp
                        150
                                           155
     Ala Ser Glu Val Ser Lys Asp Asp Ala Gly Pro Gly Ser Glu Glu
                                       170
     Lys Ser Glu Ala
30
                 180
      (2) INFORMATION FOR SEQ ID NO:144:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 153 amino acids
               (B) TYPE: amino acid
35
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
40
               (B) LOCATION: 1..153
               (D) OTHER INFORMATION: / Ceres Seq. ID 1028610
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:144:
      Met Thr Asp Pro Thr Ser Gln Glu Ala Ala Ser Lys Asp Thr Asp Leu
                                        10
45
      Thr Lys Glu Ala Glu Ser Glu Lys Lys Pro Gly Gly Ile Ser Leu Arg
                                    25
      Ile Trp Pro Pro Thr Gln Lys Thr Arg Asp Ala Val Leu Asn Arg Leu
                                40
      Ile Glu Thr Leu Ser Thr Glu Ser Ile Leu Ser Lys Arg Tyr Gly Thr
50
      Leu Lys Ser Asp Asp Ala Thr Thr Val Ala Lys Leu Ile Glu Glu Glu
                                            75
      Ala Tyr Gly Val Ala Ser Asn Ala Val Ser Ser Asp Asp Gly Ile
                                        90
55
      Lys Ile Leu Glu Leu Tyr Ser Lys Glu Ile Ser Lys Arg Met Leu Glu
                 100
                                    105
                                                       110
      Ser Val Lys Ala Arg Ser Asn Ala Ser Val Gly Asn Gly Ser Val Glu
                               120
             115
                                         125
      Asp Ala Asn Thr Asp Ala Ser Glu Val Ser Lys Asp Asp Ala Gly Pro
                  135
60
      Gly Ser Glu Glu Glu Lys Ser Glu Ala
                         150
      145
```

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(2) INFORMATION FOR SEO ID NO:145:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 94 amino acids
                (B) TYPE: amino acid
5
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
10
                (B) LOCATION: 1..94
                (D) OTHER INFORMATION: / Ceres Seq. ID 1028611
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:145:
      Met Gln Pro Pro Ser Arg Asn Leu Ser Lys Lys Arg Leu Met Val Leu
                      5
                                          10
15
      Leu Arg Met Leu Cys Arg Val Met Met Met Gly Leu Arg Phe Leu Ser
                                      25
      Phe Ile Leu Lys Arg Leu Val Arg Gly Cys Leu Asn Leu Leu Arg Leu
                                  40
      Asp Leu Met Leu Val Leu Glu Met Glu Val Trp Arg Met Leu Ile Leu
20
      Met Leu Val Arg Phe Leu Lys Met Met Leu Val Leu Val Gln Arg Arg
                                               75
      Arg Arg Val Arg Leu Glu Lys Phe Gly Phe Thr Ser Leu Leu
                      85
25
      (2) INFORMATION FOR SEQ ID NO:146:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 673 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..673
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1030069
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:146:
      atccaaaaaa aacaaaacaa aaaattatat tcaaqagaaa aaggaaaaaa tgaatttcat
                                                                              60
      ctccqatcaq qtaaaqaaac tctcaaqctc aacaccaqaq qaqccaqacc acaacaaqcc
                                                                             120
      agtegaagga accqaaacag ctacaagacc agetaccaac geegagetea tggcaagtge
                                                                             180
40
      caaggttgta gctgaagctg ctcaagccgc agctcgtaac gaatcagaca aactcgacaa
                                                                             240
      gggtaaagtc gccggagcct ctgctgatat cttagacgct tccgagaaat acggtaagtt
                                                                             300
      cgatgaaaag agtagcactg gtcactacct cgacaaggct gagaagtatc tcaacgacta
                                                                             360
      cgagtcgtca cactccaccg gtgctggtgg tcctcctcct ccgacgagtc aggctgagcc
                                                                             420
      agcaagtcag cctgagccgg cggctaagaa agacgatgaa gagtctggtg gtgggcttgg
                                                                             480
45
      aggttatgcc aagatggctc aaggtttctt gaagtgattt gatctttaat tgttgttcat
                                                                             540
      cattttcgta ataataaatt aaataactag tatcgtttgt gactagttta tgttgcttcg
                                                                             600
      tttatgttta tggggagtga cgagtgagtg taataacttc tggtgatcat gaatctaatc
                                                                             660
      catctttgtt gtg
      (2) INFORMATION FOR SEQ ID NO:147:
50
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 171 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
55
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..171
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1030070
60
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:147:
      Ser Lys Lys Asn Lys Thr Lys Asn Tyr Ile Gln Glu Lys Lys Glu Lys
                       5
                                           10
```

393 Met Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Ser Thr Pro Glu Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala Thr 40 Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val Ala 55 Glu Ala Ala Gln Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp Lys 70 Gly Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ser Glu Lys 10 85 90 Tyr Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly His Tyr Leu Asp Lys 100 105 110 Ala Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly Ala 115 120 15 Gly Gly Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln Pro 135 Glu Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Leu Gly 145 150 Gly Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys 20 165 (2) INFORMATION FOR SEQ ID NO:148: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 155 amino acids (B) TYPE: amino acid 25 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 30 (B) LOCATION: 1..155 (D) OTHER INFORMATION: / Ceres Seq. ID 1030071 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:148: Met Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Ser Thr Pro 10 35 Glu Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala Thr 20 25 Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val Ala 40 Glu Ala Ala Gln Ala Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp Lys 40 Gly Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ser Glu Lys Tyr Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly His Tyr Leu Asp Lys 90 Ala Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly Ala 45 105 100 Gly Gly Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln Pro 115 120 Glu Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Leu Gly 50 130 135 Gly Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys 150 (2) INFORMATION FOR SEQ ID NO:149: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 115 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 60 (ix) FEATURE: (A) NAME/KEY: peptide

(B) LOCATION: 1..115

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(D) OTHER INFORMATION: / Ceres Seq. ID 1030072
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:149:
     Met Ala Ser Ala Lys Val Val Ala Glu Ala Ala Gln Ala Ala Arg
                                          10
 5
     Asn Glu Ser Asp Lys Leu Asp Lys Gly Lys Val Ala Gly Ala Ser Ala
                                      25
     Asp Ile Leu Asp Ala Ser Glu Lys Tyr Gly Lys Phe Asp Glu Lys Ser
                                  4.0
      Ser Thr Gly His Tyr Leu Asp Lys Ala Glu Lys Tyr Leu Asn Asp Tyr
10
                              55
     Glu Ser Ser His Ser Thr Gly Ala Gly Gly Pro Pro Pro Thr Ser
                          70
                                              75
      Gln Ala Glu Pro Ala Ser Gln Pro Glu Pro Ala Ala Lys Lys Asp Asp
                      8.5
                                          90
15
      Glu Glu Ser Gly Gly Gly Leu Gly Gly Tyr Ala Lys Met Ala Gln Gly
                  100
                                      105
                                                          110
      Phe Leu Lys
              115
      (2) INFORMATION FOR SEQ ID NO:150:
20
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 944 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
25
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..944
                (D) OTHER INFORMATION: / Ceres Seq. ID 1032069
30
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:150:
      attgcgaatt ttaattgatc actetetete tetatetetg tteaaacate teteceaaag
                                                                              60
      taggtagaag aagatgggga cgacgttaga tgtatcaaga gcagagctag cacttgtggt
                                                                             120
      aatgtatttg aacaaagcag aggcaagaga taagttatgc agagctatac agtatggttc
                                                                             180
      caaqttcttq aqtqqtqqac aacctqqtac tqctcaaaat qttqacaaat ctactaqctt
                                                                             240
35
      agcaagaaaa gtcttccgtc ttttcaagtt tgtgaatgac ttgcatggtc ttatcagtcc
                                                                             300
      tgtgcctaaa ggaactcctc ttcctcttgt tttacttgga aagtcgaaga acgcactttt
                                                                             360
      atctacattc ttgttcctgg atcaaattgt ctggcttggg agatcaggaa tatataagaa
                                                                             420
      caaaqaacqa qctqaqttac ttggacqtat atctctcttc tgctgqatqq qatcttctqt
                                                                             480
      ctgcacaact ttagtcgagg ttggtgagat gggaaggctt tcttcatcaa tgaagaagat
                                                                             540
40
      cgaaaaggga ctcaagaatg gaaacaagta tcaggatgag gattatcgtg ctaagctaaa
                                                                             600
      aaaatcaaac gagaggtcac ttgctttgat caaatcagct atggacattg ttgtagcagc
                                                                             660
      tggtcttctt cagttagctc caacgaagat cactcctcgt gtcaccggag cttttggatt
                                                                             720
      catcacctcc atcatttctt gttaccagtt gcttccgaca cgccccaaga tcaaaacacc
                                                                             780
      ctgaagtcaa tcgaggaagc tggtgttaag gagaaagtca tttcagaaag tttctgttct
                                                                             840
45
      tottattatt aaggaaatto actgotgtta taaaacatog tataaaatac ttatttgcag
                                                                             900
      gaattacggt tattataaat ttatgcctca gattcttgat acgc
      (2) INFORMATION FOR SEQ ID NO:151:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 236 amino acids
50
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
55
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..236
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1032070
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:151:
      Met Gly Thr Thr Leu Asp Val Ser Arg Ala Glu Leu Ala Leu Val Val
60
                      5
                                           1.0
      Met Tyr Leu Asn Lys Ala Glu Ala Arg Asp Lys Leu Cys Arg Ala Ile
                                       25
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395 Gln Tyr Gly Ser Lys Phe Leu Ser Gly Gly Gln Pro Gly Thr Ala Gln 40 Asn Val Asp Lys Ser Thr Ser Leu Ala Arg Lys Val Phe Arg Leu Phe 55 5 Lys Phe Val Asn Asp Leu His Gly Leu Ile Ser Pro Val Pro Lys Gly 70 75 Thr Pro Leu Pro Leu Val Leu Leu Gly Lys Ser Lys Asn Ala Leu Leu 85 90 Ser Thr Phe Leu Phe Leu Asp Gln Ile Val Trp Leu Gly Arg Ser Gly 10 100 105 110 Ile Tyr Lys Asn Lys Glu Arg Ala Glu Leu Leu Gly Arg Ile Ser Leu 115 120 125 Phe Cys Trp Met Gly Ser Ser Val Cys Thr Thr Leu Val Glu Val Gly 135 15 Glu Met Gly Arg Leu Ser Ser Ser Met Lys Lys Ile Glu Lys Gly Leu 150 155 Lys Asn Gly Asn Lys Tyr Gln Asp Glu Asp Tyr Arg Ala Lys Leu Lys 170 175 Lys Ser Asn Glu Arg Ser Leu Ala Leu Ile Lys Ser Ala Met Asp Ile 20 185 Val Val Ala Ala Gly Leu Leu Gln Leu Ala Pro Thr Lys Ile Thr Pro 200 205 Arg Val Thr Gly Ala Phe Gly Phe Ile Thr Ser Ile Ile Ser Cys Tyr 215 220 25 Gln Leu Leu Pro Thr Arg Pro Lys Ile Lys Thr Pro 230 (2) INFORMATION FOR SEQ ID NO:152: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 220 amino acids 30 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 35 (A) NAME/KEY: peptide (B) LOCATION: 1..220 (D) OTHER INFORMATION: / Ceres Seq. ID 1032071 (xi) SEQUENCE DESCRIPTION: SEO ID NO:152: Met Tyr Leu Asn Lys Ala Glu Ala Arg Asp Lys Leu Cys Arg Ala Ile 40 Gln Tyr Gly Ser Lys Phe Leu Ser Gly Gly Gln Pro Gly Thr Ala Gln 25 Asn Val Asp Lys Ser Thr Ser Leu Ala Arg Lys Val Phe Arg Leu Phe 40 45 Lys Phe Val Asn Asp Leu His Gly Leu Ile Ser Pro Val Pro Lys Gly Thr Pro Leu Pro Leu Val Leu Leu Gly Lys Ser Lys Asn Ala Leu Leu 75 Ser Thr Phe Leu Phe Leu Asp Gln Ile Val Trp Leu Gly Arg Ser Gly 50 90 Ile Tyr Lys Asn Lys Glu Arg Ala Glu Leu Leu Gly Arg Ile Ser Leu 100 105 110 Phe Cys Trp Met Gly Ser Ser Val Cys Thr Thr Leu Val Glu Val Gly 115 120 125 55 Glu Met Gly Arg Leu Ser Ser Met Lys Lys Ile Glu Lys Gly Leu 140 135 Lys Asn Gly Asn Lys Tyr Gln Asp Glu Asp Tyr Arg Ala Lys Leu Lys 150 155 Lys Ser Asn Glu Arg Ser Leu Ala Leu Ile Lys Ser Ala Met Asp Ile 60 165 170 175 Val Val Ala Ala Gly Leu Leu Gln Leu Ala Pro Thr Lys Ile Thr Pro 185

WO 00/40695 PCT/US00/00466 396 Arg Val Thr Gly Ala Phe Gly Phe Ile Thr Ser Ile Ile Ser Cys Tyr 200 Gln Leu Leu Pro Thr Arg Pro Lys Ile Lys Thr Pro 210 215 (2) INFORMATION FOR SEQ ID NO:153: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 769 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..769 (D) OTHER INFORMATION: / Ceres Seq. ID 1033557 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:153: ggtatcgtcc gtagacqata gaggttccac actatgtaaa ggagaccgga gaaacaaagg 60 aactgggttt ggagataaga agtgggaact tgaatggctt ccttatccac ttccgttgta 120 gcttcggctt cgtcgcgttt atggaatcct gctgcctcca acggcaagat ttgcgttcct 180 tetgettege tttetetgeg taegggttgt agaeggagtt egteetetet eaettettee 240 gcttcttcgc aattgcttca ttgctcgttt ctctcgtcgc ctgtttccct agcgtctcca 300 ttttctggtt tgtccattgc gtttgatctc agcagtcaaa ctagtggact gaatggccag 360 agacgcagag gccttgtggt tagagctgga aaagctgctc tgtgtcaaac taagagaagc 420 agatcaagaa aatctctagc taggactcat ggtttccgta gaaggatgag gaccactagc 480 ggtagagcaa ccataaaqcq tcqacqtqcc aaqqqacqtt ggaacctctq tcccaaqtcc 540 aaccctagca gcggcaaacq ggcttgaagt ttgtttctct cttcactcta tctgtaatct 600 tgtttcatgt gctttttttc caccagaaga tacaatgaga aacatggttc gttgagctct 660 tttctgcgtt ttaactctct tgcatccttg attcgtttat aaatcaatgt tcttggaatt 720 catcacatgt actactatac tacttgcttc attgaatttg ttactactc (2) INFORMATION FOR SEQ ID NO:154: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 157 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..157 (D) OTHER INFORMATION: / Ceres Seq. ID 1033558 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:154: Met Ala Ser Leu Ser Thr Ser Val Val Ala Ser Ala Ser Ser Arg Leu 5 1.0 Trp Asn Pro Ala Ala Ser Asn Gly Lys Ile Cys Val Pro Ser Ala Ser 25 Leu Ser Leu Arg Thr Gly Cys Arg Arg Ser Ser Ser Ser Leu Thr Ser 40 Ser Ala Ser Ser Gln Leu Leu His Cys Ser Phe Leu Ser Ser Pro Val 55 Ser Leu Ala Ser Pro Phe Ser Gly Leu Ser Ile Ala Phe Asp Leu Ser 70 7.5 Ser Gln Thr Ser Gly Leu Asn Gly Gln Arg Arg Gly Leu Val Val 85 90 Arg Ala Gly Lys Ala Ala Leu Cys Gln Thr Lys Arg Ser Arg Ser Arg 105 Lys Ser Leu Ala Arg Thr His Gly Phe Arg Arg Arg Met Arg Thr Thr 120

60 Leu Cys Pro Lys Ser Asn Pro Ser Ser Gly Lys Arg Ala 145 150 155

135

Ser Gly Arg Ala Thr Ile Lys Arg Arg Arg Ala Lys Gly Arg Trp Asn

(2) INFORMATION FOR SEQ ID NO:155:

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WO 00/40695

PCT/US00/00466 397 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 679 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 5 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..679 10 (D) OTHER INFORMATION: / Ceres Seq. ID 1034688 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:155: aaaaaaaaac aaaacaaaaa attatattca agagaaaaaag gaaaaaaatga atttcatctc 60 cgatcaggta aagaaactct caagctcaac accagaggag ccagaccaca acaagccagt 120 cgaaggaacc gaaacagcta caagaccagc taccaacgcc gagctcatgg caagtgccaa 180 15 ggttgtagct gaagctgctc aagccgcagc tcgtaacgaa tcagacaaac tcgacaaggg 240 taaaqtcqcc qqaqcctctq ctqatatctt aqacqctqcc qaqaaatacq qtaaqttcqa 300 tgaaaagagt agcactggtc agtacctcga caaggctgag aagtatctca acgactacga 360 gtcgtcacac tccaccggtg ctggtggtcc tcctcctcca acgagtcagg ctgagccagc 420 aagtcagcct gagccggcgg ctaagaaaga cgatgaagag tctggtggtg ggcttggagg 480 20 ttatgccaag atggctcaag gtttcttgaa gtgatttgat ctttaattgt tgttcatcat 540 tttcqtaata ataaattaaa taactaqtat cqtttqtqac taqtttatqt tqcttcqttt 600 atgtttatgg ggagtgacga gtgagtgtaa taacttctgg tgatcatgaa tctaatccat 660 ctttgttgtg attatgctc (2) INFORMATION FOR SEQ ID NO:156: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 170 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..170 (D) OTHER INFORMATION: / Ceres Seq. ID 1034689 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:156: 35 Lys Lys Asn Lys Thr Lys Asn Tyr Ile Gln Glu Lys Lys Glu Lys Met 1 10 15 Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Ser Thr Pro Glu 25 40 Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala Thr Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val Ala Glu 55 Ala Ala Gln Ala Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp Lys Gly 45 7.5 Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ala Glu Lys Tyr 90 Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly Gln Tyr Leu Asp Lys Ala 105 100 110 50 Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly Ala Gly 120 115 125 Gly Pro Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln Pro Glu 135 130 140 Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Leu Gly Gly 55 155 150

165 (2) INFORMATION FOR SEQ ID NO:157:

60

(i) SEQUENCE CHARACTERISTICS:

Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys

- (A) LENGTH: 155 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS:

```
(D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
 5
               (B) LOCATION: 1..155
               (D) OTHER INFORMATION: / Ceres Seq. ID 1034690
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:157:
     Met Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Ser Thr Pro
                                        10
10
     Glu Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala Thr
               2.0
                                    25
     Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val Ala
                                40
     Glu Ala Ala Gln Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp Lys
15
                            5.5
     Gly Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ala Glu Lys
                     70
                                            75
     Tyr Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly Gln Tyr Leu Asp Lys
                    85
                                        90
20
     Ala Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly Ala
                 100
                                    105
     Gly Gly Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln Pro
                                120
      Glu Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Gly Leu Gly
25
              135
      Gly Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys
                        150
      (2) INFORMATION FOR SEQ ID NO:158:
          (i) SEQUENCE CHARACTERISTICS:
30
               (A) LENGTH: 115 amino acids
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..115
               (D) OTHER INFORMATION: / Ceres Seq. ID 1034691
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:158:
40
      Met Ala Ser Ala Lys Val Val Ala Glu Ala Ala Gln Ala Ala Arg
              5
                                         10
      Asn Glu Ser Asp Lys Leu Asp Lys Gly Lys Val Ala Gly Ala Ser Ala
      Asp Ile Leu Asp Ala Ala Glu Lys Tyr Gly Lys Phe Asp Glu Lys Ser
45
      Ser Thr Gly Gln Tyr Leu Asp Lys Ala Glu Lys Tyr Leu Asn Asp Tyr
      Glu Ser Ser His Ser Thr Gly Ala Gly Gly Pro Pro Pro Thr Ser
                                             75
50
      Gln Ala Glu Pro Ala Ser Gln Pro Glu Pro Ala Ala Lys Lys Asp Asp
                                         90
      Glu Glu Ser Gly Gly Gly Leu Gly Gly Tyr Ala Lys Met Ala Gln Gly
                                     105
      Phe Leu Lys
55
              115
      (2) INFORMATION FOR SEQ ID NO:159:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 471 base pairs
                (B) TYPE: nucleic acid
60
                (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
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(ix) FEATURE:
               (A) NAME/KEY: -
                (B) LOCATION: 1..471
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035033
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:159:
      atataaaaaa aaaacttatt ttgttcattc ttttaagttt aaaaacagtc aattcattta
     tatcatctat gatgaagaaa ctcattcaag tctcgttcac tgttatgatt atcttcacca
                                                                            120
      ttcttgtgct aggagtggtg gcaaatgagg ggctaggaaa accaaaaaaa caatgtaatg
                                                                            180
      agattctaaa gcaatccaac tgtgttgctg cagagtgtga ctctatgtgt gtgaagaaga
10
      gggggaaagg agccggttac tgctctcctt ctaaaaagtg ttactgctat tatcattgcc
      cttaacgatt ttattattta caataataag agatttatga taaagtagta aaataatgaa
      tcgggtgttt atagcctaac gattgatata gttgtctcta tcaaataaac tatatagact
                                                                            420
      aaaagtaata ataataatga caatcataac tcagaagttt ttgtttagtc c
      (2) INFORMATION FOR SEQ ID NO:160:
15
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 100 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..100
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035034
25
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:160:
      Ile Lys Lys Lys Leu Ile Leu Phe Ile Leu Leu Ser Leu Lys Thr Val
                     5
                                         10
      Asn Ser Phe Ile Ser Ser Met Met Lys Lys Leu Ile Gln Val Ser Phe
                                      25
30
      Thr Val Met Ile Ile Phe Thr Ile Leu Val Leu Gly Val Val Ala Asn
      Glu Gly Leu Gly Lys Pro Lys Lys Gln Cys Asn Glu Ile Leu Lys Gln
                              55
      Ser Asn Cys Val Ala Ala Glu Cys Asp Ser Met Cys Val Lys Lys Arg
35
                          70
                                              75
      Gly Lys Gly Ala Gly Tyr Cys Ser Pro Ser Lys Lys Cys Tyr Cys Tyr
      Tyr His Cys Pro
                  100
40
      (2) INFORMATION FOR SEQ ID NO:161:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 78 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..78
50
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035035
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:161:
      Met Met Lys Lys Leu Ile Gln Val Ser Phe Thr Val Met Ile Ile Phe
                                          10
      Thr Ile Leu Val Leu Gly Val Val Ala Asn Glu Gly Leu Gly Lys Pro
55
      Lys Lys Gln Cys Asn Glu Ile Leu Lys Gln Ser Asn Cys Val Ala Ala
      Glu Cys Asp Ser Met Cys Val Lys Lys Arg Gly Lys Gly Ala Gly Tyr
                              55
      Cys Ser Pro Ser Lys Lys Cys Tyr Cys Tyr Tyr His Cys Pro
60
                          70
      (2) INFORMATION FOR SEQ ID NO:162:
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(i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 77 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
 5
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..77
10
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035036
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:162:
      Met Lys Lys Leu Ile Gln Val Ser Phe Thr Val Met Ile Ile Phe Thr
                                          10
      Ile Leu Val Leu Gly Val Val Ala Asn Glu Gly Leu Gly Lys Pro Lys
15
                 20
                                      25
      Lys Gln Cys Asn Glu Ile Leu Lys Gln Ser Asn Cys Val Ala Ala Glu
                                  40
      Cys Asp Ser Met Cys Val Lys Lys Arg Gly Lys Gly Ala Gly Tyr Cys
                             55
20
      Ser Pro Ser Lys Lys Cys Tyr Cys Tyr Tyr His Cys Pro
                          70
      (2) INFORMATION FOR SEQ ID NO:163:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 985 base pairs
25
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
30
                (A) NAME/KEY: -
                (B) LOCATION: 1..985
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035071
          (xi) SEQUENCE DESCRIPTION: SEQ TD NO:163:
      taacgcactc tcttctccga atctagggtt tttgtctcta gcctcctcaa gctttgcaaa
35
      caatcatggc gacccagatc agcaagaaga gaaagttcgt agccgatggt gtgttttacg
                                                                             120
      ctgaactcaa tgaggttcta accagagagc ttgctgagga tggttactct ggtgttgagg
                                                                             180
      ttcgcgtcac tcctatgcgt actgagatca tcattagagc cactcgtact caaaacgttc
                                                                             240
      tcggtgagaa gggaaggagg attagggaat tgacatctct tgtacaaaag agattcaaat
                                                                             300
      ttcctcagga cagtgttgag ctttatgctg agaaggttgc taacagaggt ctttgtgcta
                                                                             360
40
      ttgctcaggc tgagtctctt cgttacaagc ttcttggtgg tcttgctgtt cgtagggcgt
                                                                             420
      gctatggtgt tttaaggttt gttatggaga gtggagctaa aggatgcgaa gtcattgtga
                                                                             180
                                                                             540
      gtggaaaact ccgtgctgca cgtgctaagt agatgaagtt caaggatggt tacatggttt
                                                                             600
      cctcaggtca gccaaccaag gaatacattg atgctgcagt taggcatgtt cttcttagac
      agggtgtgct tggtctcaag gtgaagatca tgcttgactg ggaccccaag ggcaaacaag
                                                                             660
45
      gaccgatgac accattgcct gatgttgtga tcatccatac accgaaagaa gatgatgtat
                                                                             720
      acattgcacc tgctcaggtt gttactcaag ctgcttttgt accggaagct ccattaacca
                                                                             780
                                                                             840
      ccacagatta tccagcaatg ccagttgctt agagagcctt ttcgttttct ggtttaaaac
      ccaatgggga tggttctttt tttgggacag tactagtttt ctcatgttat tctaagttaa
                                                                             900
      tacttaaagt totggaggtt gaattacatc aaggatatga atgtatgttt ttogttatot
                                                                            960
50
      tttgtttaaa ttttatttat ctatt
      (2) INFORMATION FOR SEQ ID NO:164:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 148 amino acids
                (B) TYPE: amino acid
55
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
60
                (B) LOCATION: 1..148
                (D) OTHER INFORMATION: / Ceres Seq. ID 1035072
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:164:
```

```
401
      Met Ala Thr Gln Ile Ser Lys Lys Arg Lys Phe Val Ala Asp Gly Val
                                        1.0
      Phe Tyr Ala Glu Leu Asn Glu Val Leu Thr Arg Glu Leu Ala Glu Asp
 5
      Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile
      Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg
     Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys Arg Phe Lys Phe Pro
10
     Gln Asp Ser Val Glu Leu Tyr Ala Glu Lys Val Ala Asn Arg Gly Leu
                                       90
                    85
     Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly
                 100 105 110
15
     Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu
            115 120
      Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala
              135
                                   140
     Ala Arg Ala Lys
20
     145
      (2) INFORMATION FOR SEQ ID NO:165:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 105 amino acids
               (B) TYPE: amino acid
25
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
30
               (B) LOCATION: 1..105
               (D) OTHER INFORMATION: / Ceres Seq. ID 1035073
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:165:
     Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu
                                        10
35
     Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys
                                    25
     Arg Phe Lys Phe Pro Gln Asp Ser Val Glu Leu Tyr Ala Glu Lys Val
                                40
     Ala Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr
40
     Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu
                                            75
     Arg Phe Val Met Glu Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser
                   85
45
     Gly Lys Leu Arg Ala Ala Arg Ala Lys
                 100
     (2) INFORMATION FOR SEQ ID NO:166:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 99 amino acids
50
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
55
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..99
               (D) OTHER INFORMATION: / Ceres Seq. ID 1035074
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:166:
     Met Lys Phe Lys Asp Gly Tyr Met Val Ser Ser Gly Gln Pro Thr Lys
60
                                 10
     Glu Tyr Ile Asp Ala Ala Val Arg His Val Leu Leu Arg Gln Gly Val
                                    25
```

402 Leu Gly Leu Lys Val Lys Ile Met Leu Asp Trp Asp Pro Lys Gly Lys 40 Gln Gly Pro Met Thr Pro Leu Pro Asp Val Val Ile Ile His Thr Pro 5 Lys Glu Asp Asp Val Tyr Ile Ala Pro Ala Gln Val Val Thr Gln Ala 75 Ala Phe Val Pro Glu Ala Pro Leu Thr Thr Thr Asp Tyr Pro Ala Met Pro Val Ala 10 (2) INFORMATION FOR SEQ ID NO:167: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 759 base pairs (B) TYPE: nucleic acid 15 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -20 (B) LOCATION: 1..759 (D) OTHER INFORMATION: / Ceres Seq. ID 1376587 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:167: atccgcagaa aaccctattg ctcccctttg ctcaagctca gtgcctcttt gcagcgaaag ctcaaccaac aagacgatga ttatctcaga gaacaatcgc agagagatct gcaaatacct 120 25 tttcaaagag ggagtatgct ttgctaagaa ggatttcaat ctcccaaagc atccgttgat 180 tgatgtacca aacctgcaag tgattaagct catgcagagt ttcaagtcca aggagtacgt 240 tagggagaca tttgcgtgga tgcattatta ttggtttctg actaatgaag gaattgagtt 300 cttgagaact tatcttaacc ttccttccga tgttgtccct gctactttga agaagtctgc taagcccggt ggtcgtccct ttggtggccc acctggtgat cgccaaagag gaccacctcg 420 30 ctctgatgga gaccgtccca gatttggtga ccgtgatgga taccgtggag gcccacgagg 540 tggtgatgag aagagtggag ctccagctga tttccagccg tctttccaag gaggtggtgg taggcctggt tttggccgtg gtgcaggcgg ttacagtgca gcagcaccat ctggttcagg 600 660 gttcccttga aaaatttgtt gtcatattgc gacgatggaa ggacagtttt gttttttgtt ctagttctgt ttgtgta&tg caaatctgga atctataatc tatctattac cttcaccttg 35 gtttaaacga acaaatcctg tagtttcagg attagtgcc (2) INFORMATION FOR SEQ ID NO:168: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 202 amino acids (B) TYPE: amino acid 40 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 45 (B) LOCATION: 1..202 (D) OTHER INFORMATION: / Ceres Seq. ID 1376588 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:168: Ser Ala Glu Asn Pro Ile Ala Pro Leu Cys Ser Ser Ser Val Pro Leu 10 50 Cys Ser Glu Ser Ser Thr Asn Lys Thr Met Ile Ile Ser Glu Asn Asn 25 Arg Arg Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly Val Cys Phe Ala 40 Lys Lys Asp Phe Asn Leu Pro Lys His Pro Leu Ile Asp Val Pro Asn 55 55 Leu Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser Lys Glu Tyr Val 70 75 Arg Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe Leu Thr Asn Glu 90 60 Gly Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro Ser Asp Val Val

105

Pro Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly

```
120
            115
                                                125
     Gly Pro Pro Gly Asp Arg Gln Arg Gly Pro Pro Arg Ser Asp Gly Asp
                                140
               135
     Arg Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Gly Gly Pro Arg Gly
 5
                            155
                      150
     Gly Asp Glu Lys Ser Gly Ala Pro Ala Asp Phe Gln Pro Ser Phe Gln
                        170
     Gly Gly Gly Arg Pro Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser
               180 185
10
     Ala Ala Pro Ser Gly Ser Gly Phe Pro
           195
     (2) INFORMATION FOR SEQ ID NO:169:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 177 amino acids
15
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
20
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..177
              (D) OTHER INFORMATION: / Ceres Seq. ID 1376589
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:169:
     Met Ile Ile Ser Glu Asn Asn Arg Glu Ile Cys Lys Tyr Leu Phe
25
                                     10
     Lys Glu Gly Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Pro Lys His
                2.0
                                  25
     Pro Leu Ile Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser
                              40
30
     Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr
                        55
     Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu
                                         75
     Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys
35
                                     90
     Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Gln Arg Gly
                                105
     Pro Pro Arg Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly
                             120 125
40
     Tyr Arg Gly Gly Pro Arg Gly Gly Asp Glu Lys Ser Gly Ala Pro Ala
                         135
     Asp Phe Gln Pro Ser Phe Gln Gly Gly Gly Arg Pro Gly Phe Gly
                      150 155
     Arg Gly Ala Gly Gly Tyr Ser Ala Ala Ala Pro Ser Gly Ser Gly Phe
45
                   165 170
     (2) INFORMATION FOR SEQ ID NO:170:
         (i) SEQUENCE CHARACTERISTICS:
50
              (A) LENGTH: 132 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
55
        (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..132
              (D) OTHER INFORMATION: / Ceres Seq. ID 1376590
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:170:
60
     Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp
                       10
             5
     Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg
```

PCT/US00/00466 WO 00/40695

404

30 25 20 Thr Tyr Leu Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys 4.0 Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg 55 Gln Arg Gly Pro Pro Arg Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp

Arg Asp Gly Tyr Arg Gly Gly Pro Arg Gly Gly Asp Glu Lys Ser Gly

10 Ala Pro Ala Asp Phe Gln Pro Ser Phe Gln Gly Gly Gly Arg Pro 105

Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Ala Pro Ser Gly 115 120

Ser Gly Phe Pro

15 130

5

20

25

45

- (2) INFORMATION FOR SEQ ID NO:171:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 645 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (ix) FEATURE:
    - (A) NAME/KEY: -
- (B) LOCATION: 1..645
  - (D) OTHER INFORMATION: / Ceres Seq. ID 1378581
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:171:

gatttcccat tttggttgca gcgatcggag agctctgacg gcaccagcca cctctgcggc ttcgcaaatc caaataagct gtggatgaaa cgaagaatcc tagatttctg aatggcgaca 30 ccatttatag cggggttgc agtagctgca acagcacttg ctggtagata tggaatccaa gcatggcaag cattcaaggc aaggccacca aggcccaaaa ttaagaaatt ttatgagggc ggtttccagc ctacgatgac gaaaagggaa gctgctctca ttcttggcgt cagggagagt 300 gtagcggcgg aaaaggtgaa ggaagcacac aggaaggtaa tgç agcaaa ccatccagat gcaggtggta gccatttcct agcctctaag atcaatgaag ctaaajacgt gatgctaggc 420 35 aaaactaaaa acagcggatc cgctttttga tgcaaatttc aatcaaaaga cttccttatg aatcatctta tgttatagga ctgtaatgtc aaaatgcaga ttttatgtta atttataagc 540 ttgtggtctg tttgtgacac ataaagactc gaatcttttc tttcacttgt tgtcgctgaa

- (2) INFORMATION FOR SEQ ID NO:172:
- 40 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 112 amino acids

aagaacacaa aaataaaaac gcaaaqcaga agggcgttct agtcg

- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: peptide
    - (B) LOCATION: 1..112
    - (D) OTHER INFORMATION: / Ceres Seq. ID 1378582
- 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:172:

Met Ala Thr Pro Phe Ile Ala Gly Val Ala Val Ala Ala Thr Ala Leu 10

Ala Gly Arg Tyr Gly Ile Gln Ala Trp Gln Ala Phe Lys Ala Arg Pro 25

55 Pro Arg Pro Lys Ile Lys Lys Phe Tyr Glu Gly Phe Gln Pro Thr 40

Met Thr Lys Arg Glu Ala Ala Leu Ile Leu Gly Val Arg Glu Ser Val 55 Ala Ala Glu Lys Val Lys Glu Ala His Arg Lys Val Met Val Ala Asn

60 75 70 His Pro Asp Ala Gly Gly Ser His Phe Leu Ala Ser Lys Ile Asn Glu 90

```
Ala Lys Asp Val Met Leu Gly Lys Thr Lys Asn Ser Gly Ser Ala Phe
                  100
                                      105
      (2) INFORMATION FOR SEQ ID NO:173:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 64 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
10
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..64
                (D) OTHER INFORMATION: / Ceres Seq. ID 1378583
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:173:
15
      Met Thr Lys Arg Glu Ala Ala Leu Ile Leu Gly Val Arg Glu Ser Val
                                          10
     Ala Ala Glu Lys Val Lys Glu Ala His Arg Lys Val Met Val Ala Asn
                                      25
      His Pro Asp Ala Gly Gly Ser His Phe Leu Ala Ser Lys Ile Asn Glu
20
                                  40
      Ala Lys Asp Val Met Leu Gly Lys Thr Lys Asn Ser Gly Ser Ala Phe
                              55
      (2) INFORMATION FOR SEQ ID NO:174:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 803 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
30
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..803
                (D) OTHER INFORMATION: / Ceres Seq. ID 1383462
          (xi) SEQUENCE DESCRIPTION: SEQ 1D NO:174:
35
                                                                              60
      aggacacgtt tctatctaga gagaacacaa cacatcgaac catggagaaa tcaacaagaa
                                                                             120
      ccctattcat aaccatcgtg ataacctcca tgttgctagg gtttggaaac tctgatcttg
      ctcaggacag agaggagtgt acgaaccagc tcatagaact atccacgtgt attccgtacg
                                                                             180
                                                                             240
      ttggaggaga cgccaaggct ccaacaaaag attgttgtgc agggtttggc caagttataa
      gaaagaqtqa qaaqtqttt tqcatattqq tcaqaqacaa aqatqatcct caacttqgta
                                                                             300
40
      tcaaqattaa eqcaacccta qeeqetcate tteecteeqe ttqtcatate acqqetecta
                                                                             360
      acatcactga ctgtatttcg attctgcatc tacctcgtaa ctcaacattg gctaaagagt
                                                                             420
      ttgagaactt aggaaggatt gaagacaact acaactccac atctcctaca caaattcata
                                                                             480
      aagatgggac aggaggaga aaagccgaac cagtgaagag taatggatgg aaggagaaga
                                                                             540
      gttggttggg tgttgagctt ttaatatatt tgctagtttc tctcatcttc ttttaattac
                                                                             600
45
      ttcctttctt taatqttcct ttqctttatt attacaataa cttataaqat atctatttat
                                                                             720
      atatgagaca catgcatgtg tttgcgatct atatgcatgt aaatgagtca ctggttaatt
      tttaatcacq aaqcqtacat tttaatttcq ttqttqccaa ttqatcttqt acgttagtga
      attttatgga ttatcgactt act
      (2) INFORMATION FOR SEQ ID NO:175:
50
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 197 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
55
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..197
                (D) OTHER INFORMATION: / Ceres Seq. ID 1383463
60
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:175:
      Asp Thr Phe Leu Ser Arg Glu Asn Thr Thr His Arg Thr Met Glu Lys
                                           10
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406 Ser Thr Arg Thr Leu Phe Ile Thr Ile Val Ile Thr Ser Met Leu Leu 25 Gly Phe Gly Asn Ser Asp Leu Ala Gln Asp Arg Glu Glu Cys Thr Asn 40 5 Gln Leu Ile Glu Leu Ser Thr Cys Ile Pro Tyr Val Gly Gly Asp Ala Lys Ala Pro Thr Lys Asp Cys Cys Ala Gly Phe Gly Gln Val Ile Arg Lys Ser Glu Lys Cys Val Cys Ile Leu Val Arg Asp Lys Asp Pro 10 Gln Leu Gly Ile Lys Ile Ası Ala Thr Leu Ala Ala His Leu Pro Ser 105 Ala Cys His Ile Thr Ala Pro Asn Ile Thr Asp Cys Ile Ser Ile Leu 120 His Leu Pro Arg Asn Ser Thr Leu Ala Lys Glu Phe Glu Asn Leu Gly 15 135 140 Arg Ile Glu Asp Asn Tyr Asn Ser Thr Ser Pro Thr Gln Ile His Lys 155 160 150 Asp Gly Thr Gly Gly Gly Lys Ala Glu Pro Val Lys Ser Asn Gly Trp 20 165 170 Lys Glu Lys Ser Trp Leu Gly Val Glu Leu Leu Ile Tyr Leu Leu Val 180 185 Ser Leu Ile Phe Phe 195 25 (2) INFORMATION FOR SEQ ID NO:176: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 184 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptice (B) LOCATION: 1..184 35 (D) OTHER INFORMATION: / Ceres Seq. ID 1383464 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:176: Met Glu Lys Ser Thr Arg Thr Leu Phe Ile Thr Ile Val Ile Thr Ser 10 Met Leu Gly Phe Gly Asn Ser Asp Leu Ala Gln Asp Arg Glu Glu 40 20 25 Cys Thr Asn Gln Leu Ile Glu Leu Ser Thr Cys Ile Pro Tyr Val Gly 40 Gly Asp Ala Lys Ala Pro Thr Lys Asp Cys Cys Ala Gly Phe Gly Gln 55 45 Val Ile Arg Lys Ser Glu Lys Cys Val Cys Ile Leu Val Arg Asp Lys 70 75 Asp Asp Pro Gln Leu Gly Ile Lys Ile Asn Ala Thr Leu Ala Ala His 85 90 Leu Pro Ser Ala Cys His Ile Thr Ala Pro Asn Ile Thr Asp Cys Ile 50 105 Ser Ile Leu His Leu Pro Arg Asn Ser Thr Leu Ala Lys Glu Phe Glu 120 Asn Leu Gly Arg Ile Glu Asp Asn Tyr Asn Ser Thr Ser Pro Thr Gln 135 140 55 Ile His Lys Asp Gly Thr Gly Gly Lys Ala Glu Pro Val Lys Ser 155 Asn Gly Trp Lys Glu Lys Ser Trp Leu Gly Val Glu Leu Leu Ile Tyr 165 170 Leu Leu Val Ser Leu Ile Phe Phe 60 180 (2) INFORMATION FOR SEQ ID NO:177: (i) SEQUENCE CHARACTERISTICS:

407

(2) INFORMATION FOR SEQ ID NO:179:

```
(A) LENGTH: 168 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..168
                (D) OTHER INFORMATION: / Ceres Seq. ID 1383465
10-
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:177:
     Met Leu Leu Gly Phe Gly Asn Ser Asp Leu Ala Gln Asp Arg Glu Glu
                                          10
      Cys Thr Asn Gln Leu Ile Glu Leu Ser Thr Cys Ile Pro Tyr Val Gly
                                      25
15
      Gly Asp Ala Lys Ala Pro Thr Lys Asp Cys Cys Ala Gly Phe Gly Gln
                                  40
      Val Ile Arg Lys Ser Glu Lys Cys Val Cys Ile Leu Val Arg Asp Lys
                              55
      Asp Asp Pro Gln Leu Gly Ile Lys Ile Asn Ala Thr Leu Ala Ala His
20
                                              75
                          70
      Leu Pro Ser Ala Cys His Ile Thr Ala Pro Asn Ile Thr Asp Cys Ile
                                          90
                      8.5
      Ser Ile Leu His Leu Pro Arg Asn Ser Thr Leu Ala Lys Glu Phe Glu
                                     105
                  100
                                                          110
25
      Asn Leu Gly Arg Ile Glu Asp Asn Tyr Asn Ser Thr Ser Pro Thr Gln
                                 120
                                                      125
      Ile His Lys Asp Gly Thr Gly Gly Gly Lys Ala Glu Pro Val Lys Ser
                             135
                                                 140
      Asn Gly Trp Lys Glu Lys Ser Trp Leu Gly Val Glu Leu Leu Ile Tyr
30
                         150
                                             155
      Leu Leu Val Ser Leu Ile Phe Phe
                      165
      (2) INFORMATION FOR SEQ ID NO:178:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 968 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
40
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..968
                (D) OTHER INFORMATION: / Ceres Seq. ID 1386215
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:178:
45
      aagaggaaaa caaaactgca aaaacaaaaa taaaaaaaac atcgcacaag aaaataaaag
                                                                             60
      atttgtagaa tcaactaaga aaatggctag cactatgatg actacattgc ctcagttcaa
                                                                            120
      tggtcttcga gccaccaaaa tctctgcagc tcctgtacaa ggcctggcaa gtgttcagcc
                                                                            180
      catgagacgc aagggaaatg gagctttggg tgcaaagtgt gacttcatcg gttcatcaac
                                                                            240
      aaatctgata atggtaacgt cgacgaccct gatgttgttc gcggggagat tcggacttgc
                                                                            300
50
      gccatcagcc aataggaagg cgacagctgg acttaggttg gaggcacgtg actcaggtct
                                                                            360
      acaaacgggt gacccggccg ggttcacgct tgcggacact ttggcttgtg cctccaatct
                                                                            420
      aaccccaatt tcaccttctt ccccaaatct cgagtcaatg aagacgtccc cgatgaaacg
                                                                            480
      cctctcactc cgaaatcggc aatggctctt gtcgcagctg gaaatatctc ggtggctaat
                                                                            540
      atgaataaag cggtggctct gttaaaggaa gaagctctta agaagatcat agcagctaaa
                                                                             600
55
      acggccaagt tgagagctaa aggtgcattg acgaatcttc aggatattgt tattcgacag
                                                                             660
                                                                            720
      agtaaggtca ccggcaagag gaaggaagat gaacgctgat tttctaaatc ttgatagcaa
      ttttactggt acgtagaget tttgtgtgat atatcgette tagggtteaa ggaaagaeta
                                                                             780
      cttttttgcg gttattggcc atttatcttg gtgaatttta gttcatattc agacgtgttt
                                                                            840
      gatcaaatgt ctcccagaga aagctggaga ctttgaatta ggatacatta gggcttttat
                                                                             900
60
      tttgtacaat ggtgtgta agactcagtg acattcattc atattaatcg gttccatctt
                                                                            960
      caatcttt
```

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(i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 180 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
 5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..180
10
               (D) OTHER INFORMATION: / Ceres Seq. ID 1386216
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:179:
     Arg Gly Lys Gln Asn Cys Lys Asn Lys Asn Lys Asn Ile Ala Gln
                                        10
     Glu Asn Lys Arg Phe Val Glu Ser Thr Lys Lys Met Ala Ser Thr Met
15
                                    25
                2.0
     Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg Ala Thr Lys Ile Ser
                                4 0
     Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg Lys
                             55
20
     Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe Ile Gly Ser Ser Thr
                        70
                                            75
     Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met Leu Phe Ala Gly Arg
                                        90
                     8.5
     Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala Thr Ala Gly Leu Arg
25
                100
                                    105
     Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly Asp Pro Ala Gly Phe
                               120
                                                 125
     Thr Leu Ala Asp Thr Leu Ala Cys Ala Ser Asn Leu Thr Pro Ile Ser
                            135
30
     Pro Ser Ser Pro Asn Leu Glu Ser Met Lys Thr Ser Pro Met Lys Arg
          150 155
     Leu Ser Leu Arg Asn Arg Gln Trp Leu Leu Ser Gln Leu Glu Ile Ser
     Arg Trp Leu Ile
35
                 180
      (2) INFORMATION FOR SEQ ID NO:180:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 153 amino acids
               (B) TYPE: amino acid
40
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..153
45
               (D) OTHER INFORMATION: / Ceres Seq. ID 1386217
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:180:
     Met Ala Ser Thr Met Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg
                 5
                                     10
50
      Ala Thr Lys Ile Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln
      Pro Met Arg Arg Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe
                                 40
      Ile Gly Ser Ser Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met
55
      Leu Phe Ala Gly Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala
      Thr Ala Gly Leu Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly
                                         90
      Asp Pro Ala Gly Phe Thr Leu Ala Asp Thr Leu Ala Cys Ala Ser Asn
                                     105
      Leu Thr Pro Ile Ser Pro Ser Ser Pro Asn Leu Glu Ser Met Lys Thr
```

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120
     Ser Pro Met Lys Arg Leu Ser Leu Arg Asn Arg Gln Trp Leu Leu Ser
                135
     Gln Leu Glu Ile Ser Arg Trp Leu Ile
 5
                 150
     (2) INFORMATION FOR SEQ ID NO:181:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 149 amino acids
               (B) TYPE: amino acid
10
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
15
               (B) LOCATION: 1..149
               (D) OTHER INFORMATION: / Ceres Seq. ID 1386218
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:181:
     Met Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg Ala Thr Lys Ile
                                       10
20
     Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg
                                   25
     Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe Ile Gly Ser Ser
                               40
     Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Met Leu Phe Ala Gly
25
                         55
     Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala Thr Ala Gly Leu
                                            75
                        70
     Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly Asp Pro Ala Gly
                    85
                                       90
30
     Phe Thr Leu Ala Asp Thr Leu Ala Cys Ala Ser Asn Leu Thr Pro Ile
                                   105 110
                100
     Ser Pro Ser Ser Pro Asn Leu Glu Ser Met Lys Thr Ser Pro Met Lys
            115 120 125
     Arg Leu Ser Leu Arg Asn Arg Gln Trp Leu Leu Ser Gln Leu Glu Ile
35
                           135
     Ser Arg Trp Leu Ile
     145
      (2) INFORMATION FOR SEQ ID NO:182:
          (i) SEQUENCE CHARACTERISTICS:
40
               (A) LENGTH: 65 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
45
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..65
               (D) OTHER INFORMATION: / Ceres Seq. ID 2025156
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:182:
50
     Met Ala Leu Val Ala Ala Gly Asn Ile Ser Val Ala Asn Met Asn Lys
                                        10
     Ala Val Ala Leu Leu Lys Glu Glu Ala Leu Lys Lys Ile Ile Ala Ala
      Lys Thr Ala Lys Leu Arg Ala Lys Gly Ala Leu Thr Asn Leu Gln Asp
55
                                40
      Ile Val Ile Arg Gln Ser Lys Val Thr Gly Lys Arg Lys Glu Asp Glu
                            55
     Arg
60
      (2) INFORMATION FOR SEQ ID NO:183:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 770 base pairs
```

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410
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
 5
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..770
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388499
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:183:
10
      aaaaaactca ttctcataac aattccctta actctatctc taggtgactt aagagtaaaa
                                                                             60
      aatggtagag gcagaggttg aagtagatgt ggaaatcaaa tctacggctg ataagttttt
                                                                            120
      catgttctct agaaqatcac aacatgcctc caaagccact cgctatgtac agggatgtga
                                                                            180
      tctgctcgaa ggcgagtggg gcgaagttgg aagcatactc ttgtggaaat taactgttga
                                                                            240
      tggagagcca aaagtgtcaa aggatatgat tgaggcgatt gatatgaaga tgaatatgat
15
      ccagtggagg gtgttagagg gacctctgaa ggaagagtac aatattttct cgaaaacgat
      gaaagtaagc cctaagcaag gagggtctgg aagtgtggtg aagtggaact tgaagtatga
                                                                            420
      gagaattgat gaaaaggtgg ctcacctaga gaggctcctt cagttcttcg tcgaatgtgt
      taacgagatt gaccaatacc tattgtctga gggttagggg acttttctcc ctggcttatg
                                                                            540
      tgtgggagag ttattgtatg cgtacgtgta aggactette ttgtgtttta tatgtettte
                                                                            600
20
      tcaataatgt gatatcttga gatttagaag agaggtatcc aagtgtgcta tacctcatgt
                                                                            660
      cttcaagttg gaagttgtat aacattttaa gttgcttatg agtttctcat atatgtagtg
      tgtttgttgg ttgaaaaatc atattataat gcaacgatac catgttttt
      (2) INFORMATION FOR SEQ ID NO:184:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 151 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..151
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388500
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:184:
35
      Met Val Glu Ala Glu Val Glu Val Asp Val Glu Ile Lys Ser Thr Ala
                                          10
      Asp Lys Phe Phe Met Phe Ser Arg Arg Ser Gln His Ala Ser Lys Ala
                                      25
      Thr Arg Tyr Val Gln Gly Cys Asp Leu Leu Glu Gly Glu Trp Gly Glu
40
                                  40
      Val Gly Ser Ile Leu Leu Trp Lys Leu Thr Val Asp Gly Glu Pro Lys
                              55
      Val Ser Lys Asp Met Ile Glu Ala Ile Asp Met Lys Met Asn Met Ile
                          70
45
      Gln Trp Arg Val Leu Glu Gly Pro Leu Lys Glu Glu Tyr Asn Ile Phe
                                          90
      Ser Lys Thr Met Lys Val Ser Pro Lys Gln Gly Gly Ser Gly Ser Val
                                      105
                  100
      Val Lys Trp Asn Leu Lys Tyr Glu Arg Ile Asp Glu Lys Val Ala His
50
                                  120
              115
      Leu Glu Arg Leu Gln Phe Phe Val Glu Cys Val Asn Glu Ile Asp
                              135
      Gln Tyr Leu Leu Ser Glu Gly
                          150
55
      (2) INFORMATION FOR SEQ ID NO:185:
           (i) SEQUENCE CHARACTERISTICS:
```

- (A) LENGTH: 131 amino acids

  - (B) TYPE: amino acid
- (C) STRANDEDNESS: (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
    - (ix) FEATURE:

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411
            (A) NAME/KEY: peptide
                (B) LOCATION: 1..131
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388501
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:185:
 5
     Met Phe Ser Arg Arg Ser Gln His Ala Ser Lys Ala Thr Arg Tyr Val
                                         10
     Gln Gly Cys Asp Leu Leu Glu Gly Glu Trp Gly Glu Val Gly Ser Ile
                20
                                      25
      Leu Leu Trp Lys Leu Thr Val Asp Gly Glu Pro Lys Val Ser Lys Asp
10
                                 40
     Met Ile Glu Ala Ile Asp Met Lys Met Asn Met Ile Gln Trp Arg Val
                              55
      Leu Glu Gly Pro Leu Lys Glu Glu Tyr Asn Ile Phe Ser Lys Thr Met
                          70
                                              75
15
      Lys Val Ser Pro Lys Gln Gly Gly Ser Gly Ser Val Val Lys Trp Asn
                                          90
      Leu Lys Tyr Glu Arg Ile Asp Glu Lys Val Ala His Leu Glu Arg Leu
                                     105
      Leu Gln Phe Phe Val Glu Cys Val Asn Glu Ile Asp Gln Tyr Leu Leu
20
                                 120
      Ser Glu Gly
          130
      (2) INFORMATION FOR SEQ ID NO:186:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 83 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..83
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388502
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:186:
35
      Met Ile Glu Ala Ile Asp Met Lys Met Asn Met Ile Gln Trp Arg Val
                                          10
      Leu Glu Gly Pro Leu Lys Glu Glu Tyr Asn Ile Phe Ser Lys Thr Met
                                      25
      Lys Val Ser Pro Lys Gln Gly Gly Ser Gly Ser Val Val Lys Trp Asn
40
      Leu Lys Tyr Glu Arg Ile Asp Glu Lys Val Ala His Leu Glu Arg Leu
                              55
      Leu Gln Phe Phe Val Glu Cys Val Asn Glu Ile Asp Gln Tyr Leu Leu
                                              75
45
      Ser Glu Gly
      (2) INFORMATION FOR SEQ ID NO:187:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 752 base pairs
50
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
55
                (A) NAME/KEY: -
                (B) LOCATION: 1..752
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388519
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:187:
      aatotcacgo gotttoatoa tottottoot agattogoto otocactatt ogaaaaaatg
                                                                             60
60
      gcgaaactqq tqatgttqtt qqttctctqt atcttaccqq cqatagccat ggcggcaaqq
                                                                             120
```

aggggaaata ttggaaagaa tacaatggtg gttcaaggta gcacctactg tgacatttgc

aaattegget tegagactee tgaateetee tactteatee eeggtgeaac ggtgaageta

180

412 tcatgcaaag acaggaagac aatggaagag gtttacacag acaaagctgt atcagacaaa 300 gaaggaaagt ataagttcat tgtccacgac gatcacagag accagatgtg cgatgttttg 360 cttgtgaaaa gctcggataa aacctgctct aaaatctccg ttggacgtga gaagtctcgt 420 gtgatcttga accattacag tggcattgcc tcgcagatca gacatgctaa caacatggga 480 5 tttgagaaag aagtgagtga tgtgttctgc tctgctttgt ttcagaagta tatggttgat 540 gaagatgagg atgatattaa aaaccatctc taatctctct gtttaatctt atgatctgct 600 gttttcttca ttaatgagtt tccagttatg gaagagatat atttgtattt gtttgattac 660 ttatttgttg tctttagaga tgttgactct ggtgatcgga taactatctg tttgtgtaag 720 cttcttatat attgatgtgt catttccttg cc 10 (2) INFORMATION FOR SEQ ID NO:188: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 190 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 15 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..190 20 (D) OTHER INFORMATION: / Ceres Seq. ID 1388520 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:188: Asn Leu Thr Arg Phe His His Leu Leu Pro Arg Phe Ala Pro Pro Leu 10 Phe Glu Lys Met Ala Lys Leu Val Met Leu Leu Val Leu Cys Ile Leu 25 25 Pro Ala Ile Ala Met Ala Ala Arg Arg Gly Asn Ile Gly Lys Asn Thr 40 Met Val Val Gln Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe 55 30 Glu Thr Pro Glu Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu 75 70 Ser Cys Lys Asp Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala 8.5 90 Val Ser Asp Lys Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His 35 100 105 Arg Asp Gln Met Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr 120 Cys Ser Lys Ile Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn 135 140 40 His Tyr Ser Gly Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly 155 Phe Glu Lys Glu Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys 170 Tyr Met Val Asp Glu Asp Glu Asp Ile Lys Asn His Leu 45 180 185 (2) INFORMATION FOR SEQ ID NO:189: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 171 amino acids (B) TYPE: amino acid 50 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 55 (B) LOCATION: 1..171 (D) OTHER INFORMATION: / Ceres Seq. ID 1388521 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:189: Met Ala Lys Leu Val Met Leu Leu Val Leu Cys Ile Leu Pro Ala Ile 10 5 60 Ala Met Ala Arg Arg Gly Asn Ile Gly Lys Asn Thr Met Val Val 25

Gln Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro

```
40
     Glu Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys
                             55
                                                 60
     Asp Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp
5
                         70
                                             75
     Lys Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His Arg Asp Gln
                                         90
     Met Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys
                                     105
10
     Ile Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser
                                 120
     Gly Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys
                             135
                                                 140
     Glu Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val
15
                         150
     Asp Glu Asp Glu Asp Asp Ile Lys Asn His Leu
                     165
      (2) INFORMATION FOR SEQ ID NO:190:
          (i) SEQUENCE CHARACTERISTICS:
20
                (A) LENGTH: 166 amino acids
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
25
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..166
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388522
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:190:
30
      Met Leu Leu Val Leu Cys Ile Leu Pro Ala Ile Ala Met Ala Ala Arg
                                         10
               5
      Arg Gly Asn Ile Gly Lys Asn Thr Met Val Val Gln Gly Ser Thr Tyr
                                     25
      Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro Glu Ser Ser Tyr Phe
35
                                 40
      Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys Asp Arg Lys Thr Met
                             55
      Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp Lys Glu Gly Lys Tyr
                                          75
                         70
40
      Lys Phe Ile Val His Asp Asp His Arg Asp Gln Met Cys Asp Val Leu
                                         90
      Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys Ile Ser Val Gly Arg
                                    105
      Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser Gly Ile Ala Ser Gln
45
                                 120
      Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys Glu Val Ser Asp Val
                   135
      Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val Asp Glu Asp Glu Asp
50
      Asp Ile Lys Asn His Leu
                     165
      (2) INFORMATION FOR SEQ ID NO:191:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 658 base pairs
55
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
60
                (A) NAME/KEY: -
                (B) LOCATION: 1..658
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388563
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:191:
     aagaagtttt ctctacatcc acaaaagaaa aaaagtttcc caaaaaacaa aaacttacac
                                                                             60
     aagtttagat atggatcttg aattactaca agatttgtcc aaattcaatt tcccaacacc
                                                                             120
      catcaagate cgatccaaaa cctcaaaaga caaagaagga cgaaggtgat gacgacgaag
                                                                             180
5
      atgacctcag ctgcagcaca cccacatccc aagaacacaa gattcccgcc gtcatagact
     ctccacctcc tccqccqaqa aaaccccqqc caccaccqtc aqcaccqtcq gctacqqcqq
     ctctgatgat cagatcgtgc aagaggaagc ttttagtgtc gacttgtgag ataatcatga
                                                                            360
      atcgggaaga gattgaccgt ttcttctcct ccgtctacaa tgagacgtcg actacggcta
                                                                            420
      aacggcggag aagttaccct tattgttctc gaagatgagg cttaattcaa tatttacatt
                                                                            480
10
      tttttacagt tttactggaa atattgtgaa attaattatc tgttggtgtt cggttttaaa
                                                                            540
      tatttttaat ttagttatga atatggatgg ataattttct gcaaccgcgc atattaattt
                                                                            600
      cqcatqqaqq qqtcqatqtt qtaaattqaq taataaatqa aatttaattc caattqcc
      (2) INFORMATION FOR SEQ ID NO:192:
           (i) SEQUENCE CHARACTERISTICS:
15
                (A) LENGTH: 55 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
20
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..55
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388564
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:192:
25
      Arg Ser Phe Leu Tyr Ile His Lys Arg Lys Lys Val Ser Gln Lys Thr
                                          10
      Lys Thr Tyr Thr Ser Leu Asp Met Asp Leu Glu Leu Gln Asp Leu
                                      25
      Ser Lys Phe Asn Phe Pro Thr Pro Ile Lys Ile Arg Ser Lys Thr Ser
30
              35
                                 40
      Lys Asp Lys Glu Gly Arg Arg
          50
                              55
      (2) INFORMATION FOR SEQ ID NO:193:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 50 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..50
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388565
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:193:
45
      Met Ile Arg Ser Cys Lys Arg Lys Leu Leu Val Ser Thr Cys Glu Ile
                                          10
      Ile Met Asn Arg Glu Glu Ile Asp Arg Phe Phe Ser Ser Val Tyr Asn
                                    25
      Glu Thr Ser Thr Thr Ala Lys Arg Arg Arg Ser Tyr Pro Tyr Cys Ser
50
                                  40
      Arg Arg
      (2) INFORMATION FOR SEQ ID NO:194:
           (i) SEQUENCE CHARACTERISTICS:
55
                (A) LENGTH: 39 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
60
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..39
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(D) OTHER INFORMATION: / Ceres Seq. ID 1388566
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:194:
     Met Arg Leu Asn Ser Ile Phe Thr Phe Phe Tyr Ser Phe Thr Gly Asn
                                        10
 5
     Ile Val Lys Leu Ile Ile Cys Trp Cys Ser Val Leu Asn Ile Phe Asn
                 20
                                     25
     Leu Val Met Asn Met Asp Gly
             35
      (2) INFORMATION FOR SEQ ID NO:195:
10
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 866 base pairs
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
15
         (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..866
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388793
20
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:195:
      aacggtgtgg gctaatctca cgcgctttca tcatcttctt cctagattcg ctcctccact
     120
      gctctacttc tctctctct tagattcgaa aaaatggcga aactggtgat gttgttggtt
                                                                           180
     ctctgtatct taccggcgat agccatggcg gcaaggaggg gaaatattgg aaagaataca
                                                                           240
25
     atggtggttc aaggtagcac ctactgtgac atttgcaaat tcggcttcga gactcctgaa
                                                                           300
      tectectact teateceegg tgeaacggtg aagetateat geaaagacag gaagacaatg
                                                                           360
      gaagaggttt acacagacaa agctgtatca gacaaagaag gaaagtataa gttcattgtc
                                                                           420
      cacgacgatc acagagacca gatgtgcgat gttttgcttg tgaaaagctc ggataaaacc
                                                                           480
      tgctctaaaa tctccgttgg acgtgagaag tctcgtgtga tcttgaacca ttacagtggc
                                                                           540
30
      attgcctcgc agatcagaca tgctaacaac atgggatttg agaaagaagt gagtgatgtg
                                                                           600
      ttctgctctg ctttgtttca gaagtatatg gttgatgaag atgaggatga tattaaaaac
                                                                           660
      catctctaat ctctctgttt aatcttatga tctgctgttt tcttcattaa tgagtttcga
                                                                           720
      gttatggaag agatatattt gtatttgttt gattacttat ttgttgtctt tagagatgtt
                                                                           780
      gactctggtg atcggataac tatctgtttg tgtaagcttc ttatatattg atatgtcatt
                                                                           840
35
      tccttgctta ggaagtgtct tttatt
      (2) INFORMATION FOR SEQ ID NO:196:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 171 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..171
                (D) OTHER INFORMATION: / Ceres Seq. ID 1388794
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:196:
      Met Ala Lys Leu Val Met Leu Leu Val Leu Cys Ile Leu Pro Ala Ile
                                         10
50
      Ala Met Ala Ala Arg Arg Gly Asn Ile Gly Lys Asn Thr Met Val Val
      Gln Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro
                                  40
      Glu Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys
55
      Asp Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp
                          70
      Lys Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His Arg Asp Gln
                                      90
                      8.5
60
      Met Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys
                                  105
      Ile Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser
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416
                              120
            115
                                                 125
     Gly Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys
       130 135 140
     Glu Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val
5
          150 155
     Asp Glu Asp Glu Asp Asp Ile Lys Asn His Leu
                   165
     (2) INFORMATION FOR SEQ ID NO:197:
          (i) SEQUENCE CHARACTERISTICS:
10
               (A) LENGTH: 166 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
15
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..166
               (D) OTHER INFORMATION: / Ceres Seq. ID 1388795
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:197:
20
     Met Leu Leu Val Leu Cys Ile Leu Pro Ala Ile Ala Met Ala Ala Arg
                 5
                                10
     Arg Gly Asn Ile Gly Lys Asn Thr Met Val Val Gln Gly Ser Thr Tyr
                                   25
     Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro Glu Ser Ser Tyr Phe
25
                               40
     Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys Asp Arg Lys Thr Met
                            55
     Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp Lys Glu Gly Lys Tyr
                        70
                                           75
30
     Lys Phe Ile Val His Asp Asp His Arg Asp Gln Met Cys Asp Val Leu
     Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys Ile Ser Val Gly Arg
                                105
     Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser Gly Ile Ala Ser Gln
35
             115 120
     Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys Glu Val Ser Asp Val
             135
     Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val Asp Glu Asp Glu Asp
                    150
                                           155
40
     Asp Ile Lys Asn His Leu
                    165
      (2) INFORMATION FOR SEQ ID NO:198:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 154 amino acids
45
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
50
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..154
               (D) OTHER INFORMATION: / Ceres Seq. ID 1388796
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:198:
      Met Ala Ala Arg Arg Gly Asn Ile Gly Lys Asn Thr Met Val Val Gln
55
                                       10
      Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro Glu
                                   25
      Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys Asp
                               40
60
      Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp Lys
                            55
      Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His Arg Asp Gln Met
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417 75 70 Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys Ile 90 85 Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser Gly 5 105 110 100 Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys Glu 120 125 Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val Asp 135 10 Glu Asp Glu Asp Asp Ile Lys Asn His Leu 150 (2) INFORMATION FOR SEQ ID NO:199: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 533 base pairs 15 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 20 (A) NAME/KEY: -(B) LOCATION: 1..533 (D) OTHER INFORMATION: / Ceres Seq. ID 1392041 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:199: aaccataaca caaatacacc aaaaaaaaaa aaacagaaag taacatgaga tctctcttat 60 25 tagccgtgtg cctggttctt gctttacact gcggtgaagc agccgtgtct tgcaacacgg 120 tgattqcqqa tctttaccct tqcttatcct acqtqactca qqqcqqaccq qtcccaaccc 180 tctgctgcaa cggtctcaca acactcaaga gtcaggctca aacttctgtg gaccgtcagg 240 gggtctgtcg ttgcatcaaa tctgctattg gaggactcac tctctctcct agaaccatcc 300 aaaatgcttt ggaattgcct tctaaatgtg gtgtcgatct cccttacaag ttcagccctt 360 30 ccactgactg cgacagtatc cagtgagaca agcagaaaat cttaaaggaa gctactacaa 420 gaactataat aacctaataa ttaataaatg agggcattgg tttgctagtt gctaattgat 480 cagtgatgta ttgtcatttt gaatgttcta atatcagcag gcacttatct ctg (2) INFORMATION FOR SEQ ID NO:200: (i) SEQUENCE CHARACTERISTICS: 35 (A) LENGTH: 113 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 40 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..113 (D) OTHER INFORMATION: / Ceres Seq. ID 1392042 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:200: 45 Met Arg Ser Leu Leu Ala Val Cys Leu Val Leu Ala Leu His Cys 10 Gly Glu Ala Ala Val Ser Cys Asn Thr Val Ile Ala Asp Leu Tyr Pro 25 30 Cys Leu Ser Tyr Val Thr Gln Gly Gly Pro Val Pro Thr Leu Cys Cys 50 40 Asn Gly Leu Thr Thr Leu Lys Ser Gln Ala Gln Thr Ser Val Asp Arg 55 Gln Gly Val Cys Arg Cys Ile Lys Ser Ala Ile Gly Gly Leu Thr Leu 70 75 55 Ser Pro Arg Thr Ile Gln Asn Ala Leu Glu Leu Pro Ser Lys Cys Gly 90 Val Asp Leu Pro Tyr Lys Phe Ser Pro Ser Thr Asp Cys Asp Ser Ile 105 Gln 60 (2) INFORMATION FOR SEQ ID NO:201:

(i) SEQUENCE CHARACTERISTICS:

418

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(A) LENGTH: 1028 base pairs
```

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
  - (ix) FEATURE:

5

- (A) NAME/KEY: -
- (B) LOCATION: 1..1028
- (D) OTHER INFORMATION: / Ceres Seq. ID 1393342
- 10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:201: aatatgtcct taattgctct tatcggagaa ggcttactct gtttttcagg cgaactctgt 60 ttatcttctc cgagctcgta cgagaatatt cgtaaaggtt tcttgtgagc ttaaacgatg 120 agtacccttg agaccacaag agctgagctt ggtctggtag ttgtgtattt gaataaagcg 180 gaggcgagag acaagatttg tcgagctatt caatatggtt ccaagttctt gagtgatggc 240 15 caacctggca ctgctcaaaa tgtcgacaag aacaccagct tggctaggaa agttttccgt 300 ctcttcaagt ttgtaaatga tcttcatgct ctcattagcc ctgttcccaa agggactcca 360 ctcccgcttg ttctgctcgg aaagtctaaa aacgcgttgc tgtcaacgtt cttgttccta 420 gatcaaattg tgtggcttgg caggactggg atttacaagg acaaagaacg tgctgagatt 480 cttggacgta tatcactttt ctgttggatg ggttcttcgg tttgcacatc cttggttgag 540 20 gttggggagc ttggtaggct gtcagcatca attaagaagt tagaaaaaga gatagggaac 600 aaggataaac accagaatga gcaataccgt gcaaaagtag agaaatcaaa cgagaggtca 660 ttggctctga tcaaagcagg gatggatgta gttgttgctt tcggattgct tcaattggct 720 ccaaagaaag tcactccccg agtcacaggt gctttcggat tcgcctcatc acttatctct 780 tgttatcagc tattgccgtc acatcccaag tccaagatgg tctgataaaa gggaaacaag 840 25 gaaaggcgaa ctagtttcat tgtccacatg ttccatatat agtttgtgtc tgttttacta 900 ttacttggtg ttaaatttgt cttgcttgac tactttaatg tttgtgttag attctatggt 960 atataagata atgataaaca ttgcagtctc tgtagaataa taagaacaga taagcattac 1020
  - (2) INFORMATION FOR SEQ ID NO:202:
- 30 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 235 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
  - (D) TOPOLOGY: linear
- 35 (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:

gaggtgtc

- (A) NAME/KEY: peptide
- (B) LOCATION: 1..235
- (D) OTHER INFORMATION: / Ceres Seq. ID 1393343
- 40 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:202:
  - Met Ser Thr Leu Glu Thr Thr Arg Ala Glu Leu Gly Leu Val Val 1 5 10 15
  - Tyr Leu Asn Lys Ala Glu Ala Arg Asp Lys Ile Cys Arg Ala Ile Gln 20 25 30
- 45 Tyr Gly Ser Lys Phe Leu Ser Asp Gly Gln Pro Gly Thr Ala Gln Asn 35 40 45
  - Val Asp Lys Asn Thr Ser Leu Ala Arg Lys Val Phe Arg Leu Phe Lys 50 55 60
- Phe Val Asn Asp Leu His Ala Leu Ile Ser Pro Val Pro Lys Gly Thr 50 65 70 75 80
- Pro Leu Pro Leu Val Leu Leu Gly Lys Ser Lys Asn Ala Leu Leu Ser 85 90 95
- Thr Phe Leu Phe Leu Asp Gln Ile Val Trp Leu Gly Arg Thr Gly Ile
  100 105 110
  55 Tyr Lys Asp Lys Glu Arg Ala Glu Ile Leu Gly Arg Ile Ser Leu Phe
- 115 120 125

  Cys Trp Met Gly Ser Ser Val Cys Thr Ser Leu Val Gly Val Gly Glu
- Leu Gly Arg Leu Ser Ala Ser Ile Lys Lys Leu Glu Lys Glu Ile Gly 60 145 150 150 155

140

135

60 145 150 155 160
Asn Lys Asp Lys His Gln Asn Glu Gln Tyr Arg Ala Lys Val Glu Lys
165 170 175

419 Ser Asn Glu Arg Ser Leu Ala Leu Ile Lys Ala Gly Met Asp Val Val 185 Val Ala Phe Gly Leu Leu Gln Leu Ala Pro Lys Lys Val Thr Pro Arg 200 5 Val Thr Gly Ala Phe Gly Phe Ala Ser Ser Leu Ile Ser Cys Tyr Gln 215 Leu Leu Pro Ser His Pro Lys Ser Lys Met Val 230 (2) INFORMATION FOR SEQ ID NO:203: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 653 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..653 (D) OTHER INFORMATION: / Ceres Seq. ID 1393556 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:203: tottoccaat cocaacagot toatogtttt atottocaat caaagcagot caaatcaago 60 togoaaagat gatgoacatg acottotact ggggaatoaa agocacaatt otottogatt 120 tetggaaaae tgaeteatgg ettagttaea teeteaettt aategettge ttegtettet 180 ccgctttcta tcaatacctc gagaatcgcc gcatccaatt caaatccctt tcttcctccc 240 25 gtcgtgctcc tccaccgcct cgctcttcct ccggcgtctc cgcgcctctt atccctaaat 300 ccggtaccag atccgccgct aaagctgctt cggttcttct tttcggcgtc aacgcagcga 360 teggttactt getgatgett geagetatgt ettteaaegg aggtgtttte ategegattg 420 tegteggatt aacegeegga taegetgttt ttagatetga tgaeggeggt getgataeeg 480 ccacggatga tccatgtcca tgtgcttgat aatgattgat aataaaatca aaatcggaaa 540 30 600 tagttgcttt gtaacaaaat aatccaatct aaatgtgttt cttgagtata tgc (2) INFORMATION FOR SEQ ID NO:204: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 168 amino acids 35 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 40 (A) NAME/KEY: peptide (B) LOCATION: 1..168 (D) OTHER INFORMATION: / Ceres Seq. ID 1393557 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:204: Phe Pro Ile Pro Thr Ala Ser Ser Phe Tyr Leu Pro Ile Lys Ala Ala 45 10 Gln Ile Lys Leu Ala Lys Met Met His Met Thr Phe Tyr Trp Gly Ile Lys Ala Thr Ile Leu Phe Asp Phe Trp Lys Thr Asp Ser Trp Leu Ser 50 Tyr Ile Leu Thr Leu Ile Ala Cys Phe Val Phe Ser Ala Phe Tyr Gln Tyr Leu Glu Asn Arg Arg Ile Gln Phe Lys Ser Leu Ser Ser Ser Arg Arg Ala Pro Pro Pro Pro Arg Ser Ser Gly Val Ser Ala Pro Leu 55 Ile Pro Lys Ser Gly Thr Arg Ser Ala Ala Lys Ala Ala Ser Val Leu 105 Leu Phe Gly Val Asn Ala Ala Ile Gly Tyr Leu Leu Met Leu Ala Ala 120 125 60 Met Ser Phe Asn Gly Gly Val Phe Ile Ala Ile Val Val Gly Leu Thr

135

Ala Gly Tyr Ala Val Phe Arg Ser Asp Asp Gly Gly Ala Asp Thr Ala

420 150 155 160 Thr Asp Asp Pro Cys Pro Cys Ala 165 (2) INFORMATION FOR SEQ ID NO: 205: 5 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 146 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 10 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..146 (D) OTHER INFORMATION: / Ceres Seq. ID 1393558 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:205: Met Met His Met Thr Phe Tyr Trp Gly Ile Lys Ala Thr Ile Leu Phe 5 10 Asp Phe Trp Lys Thr Asp Ser Trp Leu Ser Tyr Ile Leu Thr Leu Ile 20 20 Ala Cys Phe Val Phe Ser Ala Phe Tyr Gln Tyr Leu Glu Asn Arg Arg 40 Ile Gln Phe Lys Ser Leu Ser Ser Ser Arg Arg Ala Pro Pro Pro Arg Ser Ser Ser Gly Val Ser Ala Pro Leu Ile Pro Lys Ser Gly Thr 25 70 Arg Ser Ala Ala Lys Ala Ala Ser Val Leu Leu Phe Gly Val Asn Ala Ala Ile Gly Tyr Leu Leu Met Leu Ala Ala Met Ser Phe Asn Gly Gly 105 100 30 Val Phe Ile Ala Ile Val Val Gly Leu Thr Ala Gly Tyr Ala Val Phe 120 125 Arg Ser Asp Asp Gly Gly Ala Asp Thr Ala Thr Asp Asp Pro Cys Pro 130 135 Cys Ala 35 145 (2) INFORMATION FOR SEQ ID NO: 206: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 145 amino acids (B) TYPE: amino acid 40 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 45 (B) LOCATION: 1..145 (D) OTHER INFORMATION: / Ceres Seq. ID 1393559 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:206: Met His Met Thr Phe Tyr Trp Gly Ile Lys Ala Thr Ile Leu Phe Asp 10 50 Phe Trp Lys Thr Asp Ser Trp Leu Ser Tyr Ile Leu Thr Leu Ile Ala 25 Cys Phe Val Phe Ser Ala Phe Tyr Gln Tyr Leu Glu Asn Arg Arg Ile 40 Gln Phe Lys Ser Leu Ser Ser Ser Arg Arg Ala Pro Pro Pro Pro Arg 55 Ser Ser Ser Gly Val Ser Ala Pro Leu Ile Pro Lys Ser Gly Thr Arg 75 70 Ser Ala Ala Lys Ala Ala Ser Val Leu Leu Phe Gly Val Asn Ala Ala 90 60 Ile Gly Tyr Leu Leu Met Leu Ala Ala Met Ser Phe Asn Gly Gly Val 105 Phe Ile Ala Ile Val Val Gly Leu Thr Ala Gly Tyr Ala Val Phe Arg

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120
      Ser Asp Asp Gly Gly Ala Asp Thr Ala Thr Asp Asp Pro Cys Pro Cys
                              135
      Ala
 5
      145
      (2) INFORMATION FOR SEQ ID NO:207:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 525 base pairs
                (B) TYPE: nucleic acid
10
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
15
                (B) LOCATION: 1..525
                (D) OTHER INFORMATION: / Ceres Seq. ID 1396782
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:207:
      atgattttta cagacgagag gagacggcga caatggcgaa tttgatgatg agattaccaa
                                                                              60
      ttagcttgag aagcttctct gtttcagctt cttcatccaa cggttcgccg ccggtgatcg
20
      gaggatctag cggcggtgta ggaccgatga ttgtggaatt accgttggag aagatacgaa
                                                                             180
      gaccgttgat gcgaaccaga tccaacgatc agaacaaagt gaaagagctt atggatagta
                                                                             240
      tccgtcaaat cggtcttcaa gttccgattg atgtgattga agttgatgga acttactatg
                                                                             300
      ggttctcggg atgtcacaga tacgaggcgc atcagaagct aggtcttcca acaatacgtt
                                                                             360
      gcaaaatacg taaaggaaca aaggaaacat taaggcatca tcttcgctga agagagtttt
                                                                             420
25
      acttttatgt atattgtttg tatgatagga actcgagtat gtgtatgaaa gttaagaata
                                                                             480
      agaactttgt acaactttgt gatagattta gtgactggat tctgc
      (2) INFORMATION FOR SEQ ID NO:208:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 135 amino acids
30
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..135
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1396783
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:208:
      Asp Phe Tyr Arg Arg Glu Glu Thr Ala Thr Met Ala Asn Leu Met Met
40
                                           10
      Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser Val Ser Ala Ser Ser Ser
                                       25
      Asn Gly Ser Pro Pro Val Ile Gly Gly Ser Ser Gly Gly Val Gly Pro
                                   40
                                                       45
45
      Met Ile Val Glu Leu Pro Leu Glu Lys Ile Arg Arg Pro Leu Met Arg
      Thr Arg Ser Asn Asp Gln Asn Lys Val Lys Glu Leu Met Asp Ser Ile
                          70
                                               75
      Arg Gln Ile Gly Leu Gln Val Pro Ile Asp Val Ile Glu Val Asp Gly
50
                                           90
      Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg Tyr Glu Ala His Gln Lys
                   100
                                      105
                                                           110
      Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile Arg Lys Gly Thr Lys Glu
              115
                                  120
55
      Thr Leu Arg His His Leu Arg
                              135
          130
       (2) INFORMATION FOR SEQ ID NO:209:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 125 amino acids
60
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
```

(D) TOPOLOGY: linear

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(ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..125
 5
               (D) OTHER INFORMATION: / Ceres Seq. ID 1396784
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:209:
     Met Ala Asn Leu Met Met Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser
                                         10
     Val Ser Ala Ser Ser Ser Asn Gly Ser Pro Pro Val Ile Gly Gly Ser
10
                 20
                                      25
     Ser Gly Gly Val Gly Pro Met Ile Val Glu Leu Pro Leu Glu Lys Ile
                                 40
     Arg Arg Pro Leu Met Arg Thr Arg Ser Asn Asp Gln Asn Lys Val Lys
                             5.5
15
     Glu Leu Met Asp Ser Ile Arg Gln Ile Gly Leu Gln Val Pro Ile Asp
                          70
                                              75
     Val Ile Glu Val Asp Gly Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg
                     85
                                          90
     Tyr Glu Ala His Gln Lys Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile
20
                                     105
     Arg Lys Gly Thr Lys Glu Thr Leu Arg His His Leu Arg
      (2) INFORMATION FOR SEQ ID NO:210:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 121 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..121
                (D) OTHER INFORMATION: / Ceres Seq. ID 1396785
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:210:
35
      Met Met Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser Val Ser Ala Ser
                                          10
      Ser Ser Asn Gly Ser Pro Pro Val Ile Gly Gly Ser Ser Gly Gly Val
               20
                                      25
      Gly Pro Met Ile Val Glu Leu Pro Leu Glu Lys Ile Arg Arg Pro Leu
40
                                  40
      Met Arg Thr Arg Ser Asn Asp Gln Asn Lys Val Lys Glu Leu Met Asp
                              55
      Ser Ile Arg Gln Ile Gly Leu Gln Val Pro Ile Asp Val Ile Glu Val
                          70
45
      Asp Gly Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg Tyr Glu Ala His
                      85
                                          90
      Gln Lys Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile Arg Lys Gly Thr
                                     105
                  100
      Lys Glu Thr Leu Arg His His Leu Arg
50
              115
      (2) INFORMATION FOR SEQ ID NO:211:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 939 base pairs
                (B) TYPE: nucleic acid
55
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: DNA (genomic)
           (ix) FEATURE:
                (A) NAME/KEY: -
60
                (B) LOCATION: 1..939
                (D) OTHER INFORMATION: / Ceres Seq. ID 1396802
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:211:
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423 ataacaaacc agggtcaggg ttttatcttc catattcttt tcccactgat tcttacaatc 60 gatctaaggg aatctacqaa ctgtttqqta atttctggtg ggaaaagtct tgatcttgct 120 gtttggtttg gagtaatggt gagtgcttgg tttgttgttg gctcaccggt gatggactct 180 tegageagte catgtttatg tetggatget caeaegaegg gtaceateag gaggaagaag 240 5 atccttggca aagcaagaaa tctggaatta ggaagctctt tcactggctc acgcattgtt 300 tttaggcttt ctccgaagcg tgtatcgaga attgcgaatc gaaagagcaa aaagctcttg 360 attgtgaatg aggatgttgc tggtaattac gacgatacct tcggtgatgt acaaaagcaa 420 attgttaatt attttacata caaagctgtg aggacggttc ttcatcagct gtacgaaatg 480 aatcctcctc aatacacttg gttctataat catatcataa caaacagacc gacggatggc 540 10 aaacgtttcc tccgtgccct cggcaaggag agtcaggagc ttgcagaaag agtgatgatc 600 acgcgtcttc acttgtatgg caaatggatc aagaaatgcg accatgggaa gatataccag 660 gaaatatcgg atgaaaactt ggcgttgatg cgtgaacgcc tgatggagac cgtgatatgg 720 cetteegatg acacaaacte aaggtaatag getgaagaag acagegeaca ceacageaca 780 acatttgctt tgtcaatttg cttctcagtt tttaggactt cgagtttttt agtttgatta 840 15 900 catattacta tagtgaaaga gcagtgtgta tatgatgtct attcttggca tgtactatat gatattaggg gcaaaaaaca acacccttct tgtgctcct (2) INFORMATION FOR SEQ ID NO:212: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 248 amino acids 20 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 25 (A) NAME/KEY: peptide (B) LOCATION: 1..248 (D) OTHER INFORMATION: / Ceres Seq. ID 1396803 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:212: Ile Thr Asn Gln Gly Gln Gly Phe Ile Phe His Ile Leu Phe Pro Leu 30 10 Ile Leu Thr Ile Asp Leu Arg Glu Ser Thr Asn Cys Leu Val Ile Ser 25 Gly Gly Lys Ser Leu Asp Leu Ala Val Trp Phe Gly Val Met Val Ser 4.0 35 Ala Trp Phe Val Val Gly Ser Pro Val Met Asp Ser Ser Ser Pro 60 55 Cys Leu Cys Leu Asp Ala His Thr Thr Gly Thr Ile Arg Arg Lys Lys 75 70 Ile Leu Gly Lys Ala Arg Asn Leu Glu Leu Gly Ser Ser Phe Thr Gly 40 90 Ser Arg Ile Val Phe Arg Leu Ser Pro Lys Arg Val Ser Arg Ile Ala 105 Asn Arg Lys Ser Lys Lys Leu Leu Ile Val Asn Glu Asp Val Ala Gly 120 45 Asn Tyr Asp Asp Thr Phe Gly Asp Val Gln Lys Gln Ile Val Asn Tyr 135 Phe Thr Tyr Lys Ala Val Arg Thr Val Leu His Gln Leu Tyr Glu Met 150 155 Asn Pro Pro Gln Tyr Thr Trp Phe Tyr Asn His Ile Ile Thr Asn Arg 50 165 170 Pro Thr Asp Gly Lys Arg Phe Leu Arg Ala Leu Gly Lys Glu Ser Gln 185 Glu Leu Ala Glu Arg Val Met Ile Thr Arg Leu His Leu Tyr Gly Lys 200 55 Trp Ile Lys Lys Cys Asp His Gly Lys Ile Tyr Gln Glu Ile Ser Asp 215 220 Glu Asn Leu Ala Leu Met Arg Glu Arg Leu Met Glu Thr Val Ile Trp 230 235 Pro Ser Asp Asp Thr Asn Ser Arg 60 245 (2) INFORMATION FOR SEQ ID NO:213:

(i) SEQUENCE CHARACTERISTICS:

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424
               (A) LENGTH: 203 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
 5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..203
               (D) OTHER INFORMATION: / Ceres Seq. ID 1396804
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:213:
     Met Val Ser Ala Trp Phe Val Val Gly Ser Pro Val Met Asp Ser Ser
                                      10
     Ser Ser Pro Cys Leu Cys Leu Asp Ala His Thr Thr Gly Thr Ile Arg
                 20
                                    25
15
     Arg Lys Lys Ile Leu Gly Lys Ala Arg Asn Leu Glu Leu Gly Ser Ser
                             40
     Phe Thr Gly Ser Arg Ile Val Phe Arg Leu Ser Pro Lys Arg Val Ser
                            55
     Arg Ile Ala Asn Arg Lys Ser Lys Lys Leu Leu Ile Val Asn Glu Asp
20
                        70
                                           75
     Val Ala Gly Asn Tyr Asp Asp Thr Phe Gly Asp Val Gln Lys Gln Ile
                     85
                                        90
     Val Asn Tyr Phe Thr Tyr Lys Ala Val Arg Thr Val Leu His Gln Leu
                                    105
25
     Tyr Glu Met Asn Pro Pro Gln Tyr Thr Trp Phe Tyr Asn His Ile Ile
                                120
     Thr Asn Arg Pro Thr Asp Gly Lys Arg Phe Leu Arg Ala Leu Gly Lys
                            135
     Glu Ser Gln Glu Leu Ala Glu Arg Val Met Ile Thr Arg Leu His Leu
30
                        150
                                           155
     Tyr Gly Lys Trp Ile Lys Lys Cys Asp His Gly Lys Ile Tyr Gln Glu
                                        170
     Ile Ser Asp Glu Asn Leu Ala Leu Met Arg Glu Arg Leu Met Glu Thr
                                    185
35
     Val Ile Trp Pro Ser Asp Asp Thr Asn Ser Arg
             195
      (2) INFORMATION FOR SEQ ID NO:214:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 191 amino acids
40
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
45
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..191
               (D) OTHER INFORMATION: / Ceres Seq. ID 1396805
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:214:
      Met Asp Ser Ser Ser Pro Cys Leu Cys Leu Asp Ala His Thr Thr
50
                                         10
      Gly Thr Ile Arg Arg Lys Lys Ile Leu Gly Lys Ala Arg Asn Leu Glu
      Leu Gly Ser Ser Phe Thr Gly Ser Arg Ile Val Phe Arg Leu Ser Pro
                                 40
55
      Lys Arg Val Ser Arg Ile Ala Asn Arg Lys Ser Lys Leu Leu Ile
      Val Asn Glu Asp Val Ala Gly Asn Tyr Asp Asp Thr Phe Gly Asp Val
                                            75
      Gln Lys Gln Ile Val Asn Tyr Phe Thr Tyr Lys Ala Val Arg Thr Val
60
                    85 90
```

Leu His Gln Leu Tyr Glu Met Asn Pro Pro Gln Tyr Thr Trp Phe Tyr

105

425 Asn His Ile Ile Thr Asn Arg Pro Thr Asp Gly Lys Arg Phe Leu Arg 115 120 Ala Leu Gly Lys Glu Ser Gln Glu Leu Ala Glu Arg Val Met Ile Thr 135 140 5 Arg Leu His Leu Tyr Gly Lys Trp Ile Lys Lys Cys Asp His Gly Lys 150 155 Ile Tyr Gln Glu Ile Ser Asp Glu Asn Leu Ala Leu Met Arg Glu Arg 170 175 165 Leu Met Glu Thr Val Ile Trp Pro Ser Asp Asp Thr Asn Ser Arg 10 180 185 (2) INFORMATION FOR SEQ ID NO:215: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 753 base pairs (B) TYPE: nucleic acid 15 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -20 (B) LOCATION: 1..753 (D) OTHER INFORMATION: / Ceres Seq. ID 1397130 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:215: aaagatttgg agaaaaagtt tactaaaaaa gaatctgggg aagtaagtga gatgaaggaa 60 gaggaggaga teggaaaacc agegaaacca aaagecaaga aagatgttge teetgggagg 120 25 ttgattgata cctatgctgc acagtgcgat aactgtcaca agtggagggt gattgatagc 180 caggaggaat atgaagatat cagaagtaaa atgctcgagg atccttttaa ctgtcagaag 240 aaacagggca tgtcttgtga agagcctgct gatattgact acgattcttc tcggacttgg 300 gtcattgaca agcctggtct ccccaaaacg cctaaaggtt tcaagagaag cttagttctc 360 agaaaagatt actctaagat ggatacctac tactttactc ctaccgggaa gaagctcagg 420 30 agtogoaatg aaatogotgo ottogttgaa gooaatoogg aattoaggaa ogoacoactt 480 ggagactica atticactgt ccccaaggtc atggaagata ctgttccccc tgatccgaag 540 cttggctctc cttttccaag cactactacc actacttcag agaagagcag tgtcaagcag 600 agccataact aatatgttct cttctccggg cctttttcta ctttcttct gctccctaaa 660 gcaatgcgtt ttgagtcttt atagatttga tgttctctaa aactgttatg atattaacat 720 35 ctctaagcaa aaaaatatga tcttttgtct ctc (2) INFORMATION FOR SEQ ID NO:216: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 203 amino acids (B) TYPE: amino acid 40 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 45 (B) LOCATION: 1..203 (D) OTHER INFORMATION: / Ceres Seq. ID 1397131 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:216: Lys Asp Leu Glu Lys Lys Phe Thr Lys Lys Glu Ser Gly Glu Val Ser 5 10 50 Glu Met Lys Glu Glu Glu Glu Ile Gly Lys Pro Ala Lys Pro Lys Ala 25 20 Lys Lys Asp Val Ala Pro Gly Arg Leu Ile Asp Thr Tyr Ala Ala Gln 4.0 45 Cys Asp Asn Cys His Lys Trp Arg Val Ile Asp Ser Gln Glu Glu Tyr 55 55 Glu Asp Ile Arg Ser Lys Met Leu Glu Asp Pro Phe Asn Cys Gln Lys 70 75 Lys Gln Gly Met Ser Cys Glu Glu Pro Ala Asp Ile Asp Tyr Asp Ser 85 90 60 Ser Arg Thr Trp Val Ile Asp Lys Pro Gly Leu Pro Lys Thr Pro Lys 105 100

Gly Phe Lys Arg Ser Leu Val Leu Arg Lys Asp Tyr Ser Lys Met Asp

```
120
     Thr Tyr Tyr Phe Thr Pro Thr Gly Lys Lys Leu Arg Ser Arg Asn Glu
       130 135
                                   140
     Ile Ala Ala Phe Val Glu Ala Asn Pro Glu Phe Arg Asn Ala Pro Leu
 5
     145 150 155
     Gly Asp Phe Asn Phe Thr Val Pro Lys Val Met Glu Asp Thr Val Pro
                  165
                           170
     Pro Asp Pro Lys Leu Gly Ser Pro Phe Pro Ser Thr Thr Thr Thr
             180 185
10
     Ser Glu Lys Ser Ser Val Lys Gln Ser His Asn
            195
     (2) INFORMATION FOR SEO ID NO:217:
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 186 amino acids
15
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
20
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..186
              (D) OTHER INFORMATION: / Ceres Seq. ID 1397132
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:217:
     Met Lys Glu Glu Glu Ile Gly Lys Pro Ala Lys Pro Lys Ala Lys
25
            5
                                      10
     Lys Asp Val Ala Pro Gly Arg Leu Ile Asp Thr Tyr Ala Ala Gln Cys
               20
                                  25
     Asp Asn Cys His Lys Trp Arg Val Ile Asp Ser Gln Glu Glu Tyr Glu
                               40
30
     Asp Ile Arg Ser Lys Met Leu Glu Asp Pro Phe Asn Cys Gln Lys Lys
                           55
     Gln Gly Met Ser Cys Glu Glu Pro Ala Asp Ile Asp Tyr Asp Ser Ser
                       70
                                          75
     Arg Thr Trp Val Ile Asp Lys Pro Gly Leu Pro Lys Thr Pro Lys Gly
35
     Phe Lys Arg Ser Leu Val Leu Arg Lys Asp Tyr Ser Lys Met Asp Thr
                                  105
     Tyr Tyr Phe Thr Pro Thr Gly Lys Lys Leu Arg Ser Arg Asn Glu Ile
                              120
40
     Ala Ala Phe Val Glu Ala Asn Pro Glu Phe Arg Asn Ala Pro Leu Gly
                           135
     Asp Phe Asn Phe Thr Val Pro Lys Val Met Glu Asp Thr Val Pro Pro
                       150
                                          155
     Asp Pro Lys Leu Gly Ser Pro Phe Pro Ser Thr Thr Thr Thr Ser
4.5
                    165
                                      170
     Glu Lys Ser Ser Val Lys Gln Ser His Asn
                180
     (2) INFORMATION FOR SEQ ID NO:218:
          (i) SEQUENCE CHARACTERISTICS:
50
              (A) LENGTH: 133 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
55
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..133
              (D) OTHER INFORMATION: / Ceres Seq. ID 1397133
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:218:
60
     Met Leu Glu Asp Pro Phe Asn Cys Gln Lys Lys Gln Gly Met Ser Cys
                5
                                      10
     Glu Glu Pro Ala Asp Ile Asp Tyr Asp Ser Ser Arg Thr Trp Val Ile
```

427 25 20 30 Asp Lys Pro Gly Leu Pro Lys Thr Pro Lys Gly Phe Lys Arg Ser Leu 40 45 Val Leu Arg Lys Asp Tyr Ser Lys Met Asp Thr Tyr Tyr Phe Thr Pro 5 55 Thr Gly Lys Lys Leu Arg Ser Arg Asn Glu Ile Ala Ala Phe Val Glu Ala Asn Pro Glu Phe Arg Asn Ala Pro Leu Gly Asp Phe Asn Phe Thr 90 8.5 10 Val Pro Lys Val Met Glu Asp Thr Val Pro Pro Asp Pro Lys Leu Gly 105 100 Ser Pro Phe Pro Ser Thr Thr Thr Thr Ser Glu Lys Ser Ser Val 120 115 Lys Gln Ser His Asn 15 130 (2) INFORMATION FOR SEQ ID NO:219: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 784 base pairs (B) TYPE: nucleic acid 20 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -25 (B) LOCATION: 1..784 (D) OTHER INFORMATION: / Ceres Seq. ID 1398004 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:219: actcatcaac caaaacaaaa cataaaaaaa caaqtggaag ctttaaaacg agagggagag 60 agcaaaaatg gcgacgtcgg gaacgtacgt gacggaagtt ccgctaaaag gatcggccga 120 30 gaaatactac aagaggtgga agaacgagaa ccatgtcttc cctgatgcta tcggccacca 180 catccaaaat gttaccgttc acgaaggcga acatgactct cacgggtcta tcaggagttg 240 gaactacaca tgggatggaa aggaggaggt gttcaaggag agaagagaga tagacgatga 300 gaccaaaacg ttgacgttaa gaggacttga gggtcacgtg atggagcagc tcaaagtgta 360 cgacgtcgtc taccaattca ttcccaaatc tgaggatacc tgcatcggca aaatcacttt 420 35 aatatgggag aagcgcaacg atgattcccc agaaccaagc ggctacataa aattcgtcaa 480 gagettggtt getgacatgg gaaaccaegt tagcaaaact taatcatcat teccaeagte 540 gtcgtcgtcg tcatcatcat catcatcatc atcatcatca tcatcatcat catcatcatc 600 atcatcatca ctatctcgat ttataagtta agatgttttc agtataataa atggggtctt 660 gtggatcgtt catttctatg tgtaaaccgt ttggttctgt atgatgcttc gatatattgt 720 40 tatgttcatg atcatatgtc gggttcgata taatgattct taagattaat ttactacaca 780 tttc (2) INFORMATION FOR SEQ ID NO:220: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 151 amino acids 45 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 50 (A) NAME/KEY: peptide (B) LOCATION: 1..151 (D) OTHER INFORMATION: / Ceres Seq. ID 1398005 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:220: Met Ala Thr Ser Gly Thr Tyr Val Thr Glu Val Pro Leu Lys Gly Ser 55 10 Ala Glu Lys Tyr Tyr Lys Arg Trp Lys Asn Glu Asn His Val Phe Pro 25 Asp Ala Ile Gly His His Ile Gln Asn Val Thr Val His Glu Gly Glu 40 60 His Asp Ser His Gly Ser Ile Arg Ser Trp Asn Tyr Thr Trp Asp Gly 55

Lys Glu Glu Val Phe Lys Glu Arg Arg Glu Ile Asp Asp Glu Thr Lys

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428
                              70
                                                     75
       Thr Leu Thr Leu Arg Gly Leu Glu Gly His Val Met Glu Gln Leu Lys
      Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys Ser Glu Asp Thr Cys
 5
                    100
                                           105
       Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg Asn Asp Asp Ser Pro
                              120
                                                 125
               115
       Glu Pro Ser Gly Tyr Ile Lys Phe Val Lys Ser Leu Val Ala Asp Met
                              135
                                                         140
10
       Gly Asn His Val Ser Lys Thr
                          150
       (2) INFORMATION FOR SEQ ID NO:221:
            (i) SEQUENCE CHARACTERISTICS:
                  (A) LENGTH: 389 base pairs
15
                  (B) TYPE: nucleic acid
                  (C) STRANDEDNESS: single
                  (D) TOPOLOGY: linear
            (ii) MOLECULE TYPE: DNA (genomic)
           (ix) FEATURE:
20
                  (A) NAME/KEY: -
                  (B) LOCATION: 1..389
                  (D) OTHER INFORMATION: / Ceres Seq. ID 1399370
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:221:
      aaaaactttt toogagtogg aggtttoaga cagagaggaa gaagatgaag oggacgguga tggacaacgo gattogatoa tooggtggtgg tgttgggato attggcottt ggttacttgt 120 cactggagot tggttacaag cotttocttg aaaaggotga acaatacgaa agatototto agtottotoa acaacatoaa caacaagatg aacaagaaga agogaggtgg gacaatagoa 240 atgtogaggg gtgggaggag aaaaggtagt ottataatog gtttoaatac agacatatoa 300 cottatoaaa atttootot taatoaacog taagttoaat 360
25
30
       ctgttaacta tcaaccatgg cctgttttt
       (2) INFORMATION FOR SEQ ID NO:222:
             (i) SEQUENCE CHARACTERISTICS:
                   (A) LENGTH: 88 amino acids
                   (B) TYPE: amino acid
35
                   (C) STRANDEDNESS:
                  (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
            (ix) FEATURE:
                  (A) NAME/KEY: peptide
40
                   (B) LOCATION: 1..88
                   (D) OTHER INFORMATION: / Ceres Seq. ID 1399371
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO:222:
       Lys Leu Phe Gln Ser Arg Arg Phe Gln Thr Glu Arg Lys Lys Met Lys
                                                 10
45
      Arg Thr Val Met Asp Asn Ala Ile Arg Ser Ser Val Val Val Leu Gly
                                            25
       Ser Leu Ala Phe Gly Tyr Leu Ser Leu Glu Leu Gly Tyr Lys Pro Phe
                                        40
       Leu Glu Lys Ala Glu Gln Tyr Glu Arg Ser Leu Gln Ser Ser Gln Gln
50
                                   55
                                                          60
       His Gln Gln Gln Asp Glu Glu Glu Ala Arg Trp Asp Asn Ser Asn
       Val Glu Gly Trp Glu Glu Lys Arg
                         85
55
      (2) INFORMATION FOR SEQ ID NO:223:
             (i) SEQUENCE CHARACTERISTICS:
                   (A) LENGTH: 74 amino acids
                   (B) TYPE: amino acid
                   (C) STRANDEDNESS:
60
                  (D) TOPOLOGY: linear
            (ii) MOLECULE TYPE: peptide
            (ix) FEATURE:
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(A) NAME/KEY: peptide
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 1399372
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:223:
 5
      Met Lys Arg Thr Val Met Asp Asn Ala Ile Arg Ser Ser Val Val Val
      Leu Gly Ser Leu Ala Phe Gly Tyr Leu Ser Leu Glu Leu Gly Tyr Lys
                                      25
      Pro Phe Leu Glu Lys Ala Glu Gln Tyr Glu Arg Ser Leu Gln Ser Ser
10
                                  40
      Gln Gln His Gln Gln Gln Glu Glu Glu Glu Ala Arg Trp Asp Asn
                             5.5
      Ser Asn Val Glu Gly Trp Glu Glu Lys Arg
                         70
15
      (2) INFORMATION FOR SEQ ID NO:224:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 69 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
20
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..69
25
                (D) OTHER INFORMATION: / Ceres Seq. ID 1399373
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:224:
      Met Asp Asn Ala Ile Arg Ser Ser Val Val Leu Gly Ser Leu Ala
                                          10
      Phe Gly Tyr Leu Ser Leu Glu Leu Gly Tyr Lys Pro Phe Leu Glu Lys
30
                                      25
      Ala Glu Gln Tyr Glu Arg Ser Leu Gln Ser Ser Gln Gln His Gln Gln
                                  40
      Gln Asp Glu Gln Glu Glu Ala Arg Trp Asp Asn Ser Asn Val Glu Gly
                              55
35
      Trp Glu Glu Lys Arg
      (2) INFORMATION FOR SEQ ID NO: 225:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 745 base pairs
40
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
45
                (A) NAME/KEY: -
                (B) LOCATION: 1..745
                (D) OTHER INFORMATION: / Ceres Seq. ID 1425147
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:225:
      atttcttcgt ctctgtctcc aaaatcgaat caaaatctct aaagtttcaa tttttttgtt
                                                                             60
50
      ctgttctttt tttttttta aagaatggct tcaatttctg caactttgcc ttcgccattg
      ttactcacac agagaaaatc caatctcaca tcgattcaaa aactcccatt ttctctaact
                                                                             180
      cgaggtacga atgatctttc tccattatct cttactcgaa accctagcag catcagtctg
      atggtgaaat ctagtggaga aagctcagat tcatcgactg atctcgacgt tgttagtacg
                                                                             300
      attcagaatg tttgggataa gtctgaagat aggttaggtc ttattggttt gggttttgct
                                                                             360
55
      ggtattgtag ctctttgggc atcattgaat ctcatcacgg caattgacaa attgcccgtt
                                                                             420
      atctcgagcg gattcgaact agttggtatc ttgttctcca cgtggttcac atatcgatat
                                                                             480
      ctcttgttca aaccggacag acaggagctt tcgaaaattg tcaagaaatc agtagcggat
                                                                             540
      atacttggcc agtgaacctt gtgtgtgtga taatacttca tctttggaag atgatttgtt
                                                                             600
      tgcaagtttg taaaattaca tgacagggtg gttgttgttt ctagtccaat aatgtcatgc
                                                                             660
60
      atttgaaacc tgtaaatact ttattgttgg tttttggttg tgagcaaaat caatcttttc
                                                                            720
      taatttcaaa gattctcttt tatgt
      (2) INFORMATION FOR SEQ ID NO: 226:
```

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(i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 156 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
 5
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..156
10
                (D) OTHER INFORMATION: / Ceres Seq. ID 1425148
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:226:
     Met Ala Ser Ile Ser Ala Thr Leu Pro Ser Pro Leu Leu Leu Thr Gln
      Arg Lys Ser Asn Leu Thr Ser Ile Gln Lys Leu Pro Phe Ser Leu Thr
15
      Arg Gly Thr Asn Asp Leu Ser Pro Leu Ser Leu Thr Arg Asn Pro Ser
                                  40
                                                     45
      Ser Ile Ser Leu Met Val Lys Ser Ser Gly Glu Ser Ser Asp Ser Ser
                             55
20
      Thr Asp Leu Asp Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser
                         70
                                             75
      Glu Asp Arg Leu Gly Leu Gly Leu Gly Phe Ala Gly Ile Val Ala
                                         90
                     85
      Leu Trp Ala Ser Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val
25
                                     105
      Ile Ser Ser Gly Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe
                          120
      Thr Tyr Arg Tyr Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys
                            135
30
      Ile Val Lys Lys Ser Val Ala Asp Ile Leu Gly Gln
                         150
      (2) INFORMATION FOR SEQ ID NO:227:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 104 amino acids
35
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..104
               (D) OTHER INFORMATION: / Ceres Seq. ID 1425149
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:227:
     Met Val Lys Ser Ser Gly Glu Ser Ser Asp Ser Ser Thr Asp Leu Asp
45
                                         10
     Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser Glu Asp Arg Leu
                                     25
     Gly Leu Ile Gly Leu Gly Phe Ala Gly Ile Val Ala Leu Trp Ala Ser
                                 40
50
      Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val Ile Ser Ser Gly
      Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe Thr Tyr Arg Tyr
                                             75
      Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys Ile Val Lys Lys
55
                     85
      Ser Val Ala Asp Ile Leu Gly Gln
                 100
      (2) INFORMATION FOR SEQ ID NO:228:
           (i) SEQUENCE CHARACTERISTICS:
60
               (A) LENGTH: 769 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
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(D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
 5
                (B) LOCATION: 1..769
                (D) OTHER INFORMATION: / Ceres Seq. ID 1441102
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:228:
      ccacaacaca acttcagctt taagacagct aaaactaaat acaaagaaac agagatgggt
                                                                             60
      ttqqttacag aggaggtgag agctaaggca gagatgtaca ccggagatga gatatqtaqa
                                                                             120
10
      qaqaaqacaa agtgtttcct caaggaaata tctatqccca atggtttatt accattqaaq
                                                                             180
      gatatagaag aggttgggta tgacagagag tcaggtgttg tatggctgaa gcagaagaag
                                                                             240
      agcatcactc acaagttcac agagatagat aagcttgtct cctatggaac cgaagtcaca
                                                                             300
      qccattqttq agaccqqqaa aatcaagaag ctcactqqag tcaaaqccaa qqaacttctt
                                                                             360
      atttgggtca ctatcaatga gatctatacc gaggaacccc ctaccaagat cacgttcaag
                                                                             420
15
      acaccgacca cactgtccag gactttcccg gtcacagctt ttatagtccc agaagaacct
                                                                             480
      gccaaggagg aacctgccaa agaggaacct gccaaggaga agagcagcga agccaccgag
                                                                             540
      gccaaggagg ctgttgcaat caaggaggct gtcgcagtca aagaggcggc ctaatcagct
                                                                             600
      cagcttcagc agataaaaga gacccgaatt tttaagtgta cttcaagaat ttgaacaaat
                                                                             660
      catggagagt tagtgtttct cactgctcta actatcatgt attatctacc taccacttgt
                                                                             720
20
      tgctgcttgt tttacctgta ataagatatc aatatcatcc tgttccact
      (2) INFORMATION FOR SEQ ID NO:229:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 197 amino acids
                (B) TYPE: amino acid
25
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
30
                (B) LOCATION: 1..197
                (D) OTHER INFORMATION: / Ceres Seq. ID 1441103
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:229:
      Pro Gln His Asn Phe Ser Phe Lys Thr Ala Lys Thr Lys T, ~ Lys Glu
                     5
                                          10
35
      Thr Glu Met Gly Leu Val Thr Glu Glu Val Arg Ala Lys Ala Glu Met
                                     25
      Tyr Thr Gly Asp Glu Ile Cys Arg Glu Lys Thr Lys Cys Phe Leu Lys
                                  40
      Glu Ile Ser Met Pro Asn Gly Leu Leu Pro Leu Lys Asp Ile Glu Glu
40
      Val Gly Tyr Asp Arg Glu Ser Gly Val Val Trp Leu Lys Gln Lys Lys
                                              75
                          70
      Ser Ile Thr His Lys Phe Thr Glu Ile Asp Lys Leu Val Ser Tyr Gly
                                          90
                      85
      Thr Glu Val Thr Ala Ile Val Glu Thr Gly Lys Ile Lys Lys Leu Thr
45
                  100
                                      105
                                                          110
      Gly Val Lys Ala Lys Glu Leu Leu Ile Trp Val Thr Ile Asn Glu Ile
                                  120
                                                      125
             115
      Tyr Thr Glu Glu Pro Pro Thr Lys Ile Thr Phe Lys Thr Pro Thr Thr
50
                             135
                                                  140
      Leu Ser Arg Thr Phe Pro Val Thr Ala Phe Ile Val Pro Glu Glu Pro
                                              155
                         150
      Ala Lys Glu Glu Pro Ala Lys Glu Glu Pro Ala Lys Glu Lys Ser Ser
                                         170
                                                              175
                      165
55
      Glu Ala Thr Glu Ala Lys Glu Ala Val Ala Ile Lys Glu Ala Val Ala
                                   185
                  180
                                                          190
      Val Lys Glu Ala Ala
             195
      (2) INFORMATION FOR SEQ ID NO:230:
60
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 179 amino acids

(B) TYPE: amino acid

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432
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
 5
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..179
               (D) OTHER INFORMATION: / Ceres Seq. ID 1441104
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:230:
     Met Gly Leu Val Thr Glu Glu Val Arg Ala Lys Ala Glu Met Tyr Thr
10
                                         10
      Gly Asp Glu Ile Cys Arg Glu Lys Thr Lys Cys Phe Leu Lys Glu Ile
                                     25
      Ser Met Pro Asn Gly Leu Leu Pro Leu Lys Asp Ile Glu Glu Val Gly
                                 40
15
      Tyr Asp Arg Glu Ser Gly Val Val Trp Leu Lys Gln Lys Lys Ser Ile
                              55
      Thr His Lys Phe Thr Glu Ile Asp Lys Leu Val Ser Tyr Gly Thr Glu
                         70
                                              75
     Val Thr Ala Ile Val Glu Thr Gly Lys Ile Lys Lys Leu Thr Gly Val
20
                                         90
     Lys Ala Lys Glu Leu Leu Ile Trp Val Thr Ile Asn Glu Ile Tyr Thr
                                     105
     Glu Glu Pro Pro Thr Lys Ile Thr Phe Lys Thr Pro Thr Thr Leu Ser
                                120
                                                    125
             115
25
     Arg Thr Phe Pro Val Thr Ala Phe Ile Val Pro Glu Glu Pro Ala Lys
                            135
                                                140
      Glu Glu Pro Ala Lys Glu Glu Pro Ala Lys Glu Lys Ser Ser Glu Ala
                        150 155
      Thr Glu Ala Lys Glu Ala Val Ala Ile Lys Glu Ala Val Ala Val Lys
30
                                        170
     Glu Ala Ala
      (2) INFORMATION FOR SEQ ID NO:231:
          (i) SEQUENCE CHARACTERISTICS:
35
               (A) LENGTH: 166 amino acids
               (B) TYPE: amino acid
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..166
                (D) OTHER INFORMATION: / Ceres Seq. ID 1441105
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:231:
45
     Met Tyr Thr Gly Asp Glu Ile Cys Arg Glu Lys Thr Lys Cys Phe Leu
                                         10
     Lys Glu Ile Ser Met Pro Asn Gly Leu Leu Pro Leu Lys Asp Ile Glu
                                     25
     Glu Val Gly Tyr Asp Arg Glu Ser Gly Val Val Trp Leu Lys Gln Lys
50
                                 40
     Lys Ser Ile Thr His Lys Phe Thr Glu Ile Asp Lys Leu Val Ser Tyr
                             55
     Gly Thr Glu Val Thr Ala Ile Val Glu Thr Gly Lys Ile Lys Lys Leu
                          70
                                             75
55
     Thr Gly Val Lys Ala Lys Glu Leu Leu Ile Trp Val Thr Ile Asn Glu
                                         90
      Ile Tyr Thr Glu Glu Pro Pro Thr Lys Ile Thr Phe Lys Thr Pro Thr
```

100 105 110 Thr Leu Ser Arg Thr Phe Pro Val Thr Ala Phe Ile Val Pro Glu Glu

Pro Ala Lys Glu Glu Pro Ala Lys Glu Glu Pro Ala Lys Glu Lys Ser

120

135

115

PCT/US00/00466

WO 00/40695 433 Ser Glu Ala Thr Glu Ala Lys Glu Ala Val Ala Ile Lys Glu Ala Val 155 150 Ala Val Lys Glu Ala Ala 165 (2) INFORMATION FOR SEQ ID NO:232: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 607 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE:

- (A) NAME/KEY: -
- (B) LOCATION: 1..607
- 15 (D) OTHER INFORMATION: / Ceres Seq. ID 1447480
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:232:

aaaatctcta aagtttcaat tttttkgttc tgttcttttt ttttttaaag aatggcttca atttctgcaa ctttgccttc gccattgtta ctcacacaga gaaaatccaa tctcacatcg 120 attcaaaaac tcccattttc tctaactcga ggtacgaatg atctttctcc attatctctt 180 actcgaaacc ctagcagcat cagtctgatg gtgaaagcta gtggagaaag ctcagattca 240 togactgate togacgttgt tagtacgatt cagaatgttg caattgacaa attgcccgtt 300 atctcgagcg gattcgaact agttggtatc ttgttctcca cgtggttcac atatcgatat 360 ctcttgttca aaccggacag acaggagctt tcgaaaattg tcaagaaatc agtagcggat 420 atacttggcc agtgaacctt gtgtgtgtga taatacttca tctttggaag atgatttgtt 480 tgcaagtttg taaaattaca tgacagggtg gttgttgttt ctagtccaat aatgtcatgc 540 atttgaaacc tgtaaatact ttattgttgg tttttggttg tgagcaaaat caatcttttc 600 taatttc

- (2) INFORMATION FOR SEQ ID NO:233:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 127 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
- 35 (ix) FEATURE:

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- (A) NAME/KEY: peptide
- (B) LOCATION: 1..127
- (D) OTHER INFORMATION: / Ceres Seq. ID 1447481
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:233:

40 Met Ala Ser Ile Ser Ala Thr Leu Pro Ser Pro Leu Leu Thr Gln 19 Arg Lys Ser Asn Leu Thr Ser Ile Gln Lys Leu Pro Phe Ser Leu Thr

25

Arg Gly Thr Asn Asp Leu Ser Pro Leu Ser Leu Thr Arg Asn Pro Ser 40

Ser Ile Ser Leu Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser 55 Thr Asp Leu Asp Val Val Ser Thr Ile Gln Asn Val Ala Ile Asp Lys

70 75 Leu Pro Val Ile Ser Ser Gly Phe Glu Leu Val Gly Ile Leu Phe Ser

90 Thr Trp Phe Thr Tyr Arg Tyr Leu Leu Phe Lys Pro Asp Arg Gln Glu 105 100

Leu Ser Lys Ile Val Lys Lys Ser Val Ala Asp Ile Leu Gly Gln 120 115

- (2) INFORMATION FOR SEQ ID NO:234:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 75 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide

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(ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447482
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:234:
      Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser Thr Asp Leu Asp
                      5
                                          10
      Val Val Ser Thr Ile Gln Asn Val Ala Ile Asp Lys Leu Pro Val Ile
                                      25
10
      Ser Ser Gly Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe Thr
                                  40
      Tyr Arg Tyr Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys Ile
                              55
      Val Lys Lys Ser Val Ala Asp Ile Leu Gly Gln
15
                          70
      (2) INFORMATION FOR SEQ ID NO:235:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 668 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..668
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447577
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:235:
      aaaaaaaaac aaaacaaaaa attatattca agagaaaaaag gaaaaaaatga atttcatctc
      cgatcaggta aagaaactct caagctcaac accagaggcc agaccacaac aagccagtcg
                                                                             120
30
      aaqqaaccqa aacaqctaca aqaccaqcta ccaacqccqa gctcatqqca aqtqccaaqq
                                                                             180
      ttgtagctga agctgctcaa gccgcagctc gtaacgaatc agacaaactc gacaagggta
                                                                             240
      aagtcgccqg agcctctgct gatatct ag acgctgccga gaaatacggt aagttcgatg
                                                                             300
      aaaaqagtaq cactqqtcaq tacctcgaua aggctgagaa gtatctcaac gactacgagt
                                                                             360
      cgtcacactc caccggtgct ggtggtcctc ctcctccgac gagtcaggct gagccagcaa
                                                                             420
35
      gtcagcctga gccggcggct aagaaagacg atgaagagtc tggtggtggg cttggaggtt
                                                                             480
      atgccaagat ggctcaaggt ttcttgaagt gatttgatct ttaattgttg ttcatcattt
                                                                             540
      tcgtaataat aaattaaata actagtatcg tttgtgacta gtttatgttg cttcgtttat
                                                                             600
      gtttatgggg agtgacgagt gagtgtaata acttctggtg atcatgaatc taatccatct
                                                                             660
      ttgttgtc
40
      (2) INFORMATION FOR SEQ ID NO:236:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 62 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..62
50
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447578
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:236:
      Lys Lys Asn Lys Thr Lys Asn Tyr Ile Gln Glu Lys Lys Glu Lys Met
                                           10
      Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Thr Pro Glu
55
      Ala Arg Pro Gln Gln Ala Ser Arg Arg Asn Arg Asn Ser Tyr Lys Thr
              35
                                  40
      Ser Tyr Gln Arg Arg Ala His Gly Lys Cys Gln Gly Cys Ser
                              55
60
      (2) INFORMATION FOR SEQ ID NO:237:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 115 amino acids
```

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(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..115
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447579
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:237:
10
      Met Ala Ser Ala Lys Val Val Ala Glu Ala Ala Gln Ala Ala Arg
                      5
                                         10
      Asn Glu Ser Asp Lys Leu Asp Lys Gly Lys Val Ala Gly Ala Ser Ala
                                      25
      Asp Ile Leu Asp Ala Ala Glu Lys Tyr Gly Lys Phe Asp Glu Lys Ser
15
                                  40
      Ser Thr Gly Gln Tyr Leu Asp Lys Ala Glu Lys Tyr Leu Asn Asp Tyr
                              55
      Glu Ser Ser His Ser Thr Gly Ala Gly Gly Pro Pro Pro Pro Thr Ser
20
      Gln Ala Glu Pro Ala Ser Gln Pro Glu Pro Ala Ala Lys Lys Asp Asp
                                          90
      Glu Glu Ser Gly Gly Gly Leu Gly Gly Tyr Ala Lys Met Ala Gln Gly
                  100
                                     105
      Phe Leu Lys
25
             115
      (2) INFORMATION FOR SEQ ID NO:238:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 69 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
35
                (B) LOCATION: 1..69
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447580
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:238:
      Met Lys Arg Val Ala Leu Val Ser Thr Ser Thr Arg Leu Arg Ser Ile
                                          10
40
      Ser Thr Thr Thr Ser Arg His Thr Pro Pro Val Leu Val Val Leu Leu
                                      25
      Leu Arg Arg Val Arg Leu Ser Gln Gln Val Ser Leu Ser Arg Arg Leu
                                  40
      Arg Lys Thr Met Lys Ser Leu Val Val Gly Leu Glu Val Met Pro Arg
45
      Trp Leu Lys Val Ser
      (2) INFORMATION FOR SEQ ID NO:239:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 717 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
55
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..717
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447922
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:239:
60
      cttcgtctct gtctccaaaa tcgaatcaaa atctctaaag tttcaatttt tttgttctgt
      tctttttttt tttttaaaga atggcttcaa tttctgcaac tttgccttcg ccattgttac
```

tcacacagag aaaatccaat ctcacatcga ttcaaaaaact cccattttct ctaactcgag

60

436

```
gtacgaatga totttotoca ttatototta otogaaacco tagcagcato agtotgatgg
                                                                             240
      tgaaagctag tggagaaagc tcagattcat cgactgatct cgacgttgtt agtacgattc
                                                                             300
      agaatgtttg ggataagtct gaagataggt taggtcttat tggtttgggt tttgctggta
                                                                             360
      ttgtagctct ttgggcatca ttgaatctca tcacggcaat tgacaaattg cccgttatct
                                                                             420
 5
      cgagcggatt cgaactagtt ggtatcttgt tctccacgtg gttcacatat cgatatctct
                                                                             480
      tgttcaaacc ggacagacag gagctttcga aaattgtcaa gaaatcagta gcggatatac
                                                                             540
      ttggccagtg aaccttgtgt gtgtgataat acttcatctt tggaagatga tttgtttgca
                                                                             600
      agtttgtaaa attacatgac agggtggttg ttgtttctag tccaataatg tcatgcattt
                                                                             660
      gaaacctgta aatactttat tgttggtttt tggttgtgag caaaatcaat cttttct
10
      (2) INFORMATION FOR SEQ ID NO:240:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 156 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
15
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..156
20
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447923
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:240:
      Met Ala Ser Ile Ser Ala Thr Leu Pro Ser Pro Leu Leu Leu Thr Gln
      Arg Lys Ser Asn Leu Thr Ser Ile Gln Lys Leu Pro Phe Ser Leu Thr
25
                                      25
      Arg Gly Thr Asn Asp Leu Ser Pro Leu Ser Leu Thr Arg Asn Pro Ser
                                  40
      Ser Ile Ser Leu Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser
                              55
30
      Thr Asp Leu Asp Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser
                          70
                                              75
      Glu Asp Arg Leu Gly Leu Ile Gly Leu Gly Phe Ala Gly Ile Val Ala
                      85
                                          90
      Leu Trp Ala Ser Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val
35
                                      105
                  100
                                                           110
      Ile Ser Ser Gly Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe
                               120
                                                      125
              115
      Thr Tyr Arg Tyr Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys
                              135
                                                  140
40
      Ile Val Lys Lys Ser Val Ala Asp Ile Leu Gly Gln
                          150
      (2) INFORMATION FOR SEQ ID NO:241:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 104 amino acids
45
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
50
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..104
                (D) OTHER INFORMATION: / Ceres Seq. ID 1447924
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:241:
      Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser Thr Asp Leu Asp
55
                                          10
      Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser Glu Asp Arg Leu
                                      25
      Gly Leu Ile Gly Leu Gly Phe Ala Gly Ile Val Ala Leu Trp Ala Ser
                                  40
      Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val Ile Ser Ser Gly
60
                              55
                                                   60
```

Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe Thr Tyr Arg Tyr

437 75 70 65 80 Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys Ile Val Lys Lys 90 85 Ser Val Ala Asp Ile Leu Gly Gln 5 100 (2) INFORMATION FOR SEQ ID NO:242: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 656 base pairs (B) TYPE: nucleic acid 10 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -15 (B) LOCATION: 1..656 (D) OTHER INFORMATION: / Ceres Seq. ID 1448012 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:242: ggaatttete acteteteta tetetettag ceagetetga ceatttegat tttttteegg 60 tgaaaaggga gcagaaacat ggttgtacgt atcagattat cgagatttgg atgcaaaaat 120 20 cggccatttt ttagggttat ggctgctgat agcagatctc caagagacgg gaagcatctt 180 gaggtettag gttaetteaa teetttgeea ggeeaggaeg gtggtaagag gatgggtete 240 aagttcgatc gaattaagta ctggttatct gttggtgctc agccatcaga cccggttcaa 300 cgccttcttt tcagatccgg tttacttcct cctcctccaa tggtggctat gggacgtaaa 360 ggtggagcac gagacacacg cccagttgat ccaatgactg gtcgctatgt ggatgcagag 420 25 aataaaacag ttaatgccaa tgataaccag cctaaggaag aggattcaga agacaagatt 480 gcatgattca ttagccttct gtcgtcgtag cttttcaagt tcactttgtt gtcgattata 540 ttgtgtaatg cagcattaga caactgactt gtttcctttg tttggcgata aacggcaagg 600 tqtttqqcac tttttqcaga aacggcacat attttqcatt qqqatatttt aatttt (2) INFORMATION FOR SEQ ID NO:243: 30 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 135 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 35 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..135 (D) OTHER INFORMATION: / Ceres Seq. ID 1448013 40 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:243: Met Val Val Arg Ile Arg Leu Ser Arg Phe Gly Cys Lys Asn Arg Pro 5 10 Phe Phe Arg Val Met Ala Ala Asp Ser Arg Ser Pro Arg Asp Gly Lys 45 His Leu Glu Val Leu Gly Tyr Phe Asn Pro Leu Pro Gly Gln Asp Gly 40 Gly Lys Arg Met Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser 55 Val Gly Ala Gln Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser 50 75 70 Gly Leu Leu Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly 90 95 85 Ala Arg Asp Thr Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp 105 55 Ala Glu Asn Lys Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu 120 Asp Ser Glu Asp Lys Ile Ala 130 (2) INFORMATION FOR SEQ ID NO:244: 60 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 115 amino acids

(B) TYPE: amino acid

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(C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
 5
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..115
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448014
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:244:
      Met Ala Ala Asp Ser Arg Ser Pro Arg Asp Gly Lys His Leu Glu Val
10
                                          10
      Leu Gly Tyr Phe Asn Pro Leu Pro Gly Gln Asp Gly Gly Lys Arg Met
                                      25
      Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser Val Gly Ala Gln
                                  40
15
      Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser Gly Leu Leu Pro
                              55
      Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly Ala Arg Asp Thr
                          70
      Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp Ala Glu Asn Lys
20
                                         90
                     85
      Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu Asp Ser Glu Asp
                                     105
                 100
      Lys Ile Ala
              115
25
      (2) INFORMATION FOR SEQ ID NO:245:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 84 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..84
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448015
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:245:
      Met Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser Val Gly Ala
                                          10
      Gln Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser Gly Leu Leu
40
                 2.0
                                      25
      Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly Ala Arg Asp
                                  40
                                                      45
      Thr Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp Ala Glu Asn
                             55
45
      Lys Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu Asp Ser Glu
      65
                          70
                                              75
      Asp Lys Ile Ala
      (2) INFORMATION FOR SEQ ID NO:246:
50
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 458 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
55
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..458
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448135
60
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:246:
      aattttctag ggttttgaag agtgtctctc gtcgccgttg taattcctct gttagcaaat
```

cgacaaaatg ggtcactcta atgtatggaa ctctcatccg aagaagtacg gtcctggatc

439

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togtttatgo ogtgtgtgog ggaactogca ogggetgato oggaagtatg gtttgaactg
     ctgcagacag tgtttccgta gcaacgctaa ggagattgga ttcattaagt accgttaatc
                                                                           240
     aagcaccaac ttcatgattg atgcttaatg atataaacat gaaggcgtcg atgggtttgg
                                                                           300
     cttttaagct tttgtagttt ttgaaatttt tacttttgag aaccattgtt attttgggag
 5
     ttaattaagt tgttgaacct ctcattaagc atgtcttatt ttggattaat gatgttttgg
                                                                           420
     ctattctcgy atttttgttt tatcagtcaa atttgact
     (2) INFORMATION FOR SEQ ID NO:247:
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 51 amino acids
10
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
15
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..51
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448136
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:247:
     Phe Ser Arg Val Leu Lys Ser Val Ser Arg Arg Arg Cys Asn Ser Ser
20
                                         10
     Val Ser Lys Ser Thr Lys Trp Val Thr Leu Met Tyr Gly Thr Leu Ile
                      25
                20
     Arg Arg Ser Thr Val Leu Asp Leu Val Tyr Ala Val Cys Ala Gly Thr
                      40
25
     Arg Thr Gly
         50
      (2) INFORMATION FOR SEQ ID NO:248:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 56 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
               (A) NAME/KEY: peptide
                (B) LOCATION: 1..56
               (D) OTHER INFORMATION: / Ceres Seq. ID 1448137
          (xi) SEOUENCE DESCRIPTION: SEO ID NO:248:
     Met Gly His Ser Asn Val Trp Asn Ser His Pro Lys Lys Tyr Gly Pro
40
                                         10
     Gly Ser Arg Leu Cys Arg Val Cys Gly Asn Ser His Gly Leu Ile Arg
                                     25
     Lys Tyr Gly Leu Asn Cys Cys Arg Gln Cys Phe Arg Ser Asn Ala Lys
                                 40
            35
45
     Glu Ile Gly Phe Ile Lys Tyr Arg
                          55
      (2) INFORMATION FOR SEQ ID NO:249:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 38 amino acids
50
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
55
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..38
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448138
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:249:
     Met Glu Leu Ser Ser Glu Glu Val Arg Ser Trp Ile Ser Phe Met Pro
60
                                         10
                  5
      Cys Val Arg Glu Leu Ala Arg Ala Asp Pro Glu Val Trp Phe Glu Leu
```

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Leu Gln Thr Val Phe Pro
              35
      (2) INFORMATION FOR SEQ ID NO:250:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 603 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
10
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..603
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448185
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:250:
15
      gaagaagaga gtatcgttgt gaggtttggg gatcgcgaaa atggaggttc caggttcatc
      gaagaagatg atcgcaacgc aagaagagat gtctgcagct aaaatagcac ttggatctag
                                                                             120
      agatatgtgc gctcatctct tgattccgct caacaaatgt cgtcaggctg agttttacct
                                                                             180
      tccatggaaa tgtgaagacg agcgtcacgt ttatgagaag tgtgaatacg agcttgttat
                                                                             240
      ggaqaqaatq cttqcqatqa agaaqatccq tgaaqaaqaa gctttggcta aacagaataa
                                                                             300
20
      actacaaqqa aacqctqctq ttcctcttat ccctaaaact qctaatqctt aqqattcqat
                                                                             360
      teetteteea atttgeegat teeagattee gggattetet ggaactgtga agatggtgg
      ggctctgctt ttcaatcttt ttcttcttqc tagtgatgaa aaattgttgc tacattttca
                                                                             480
      gatgtgaatg ttcaagattc tctatctttt ttttttttt gaactctttt taacttggag
                                                                             540
      tggtttccca aaaaataaga tgcaaaactc atctttttgt tggttttcta tctttaatct
25
      ata
      (2) INFORMATION FOR SEQ ID NO:251:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 103 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
35
                (B) LOCATION: 1..103
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448186
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:251:
      Met Glu Val Pro Gly Ser Ser Lys Lys Met Ile Ala Thr Gln Glu Glu
                      5
                                          10
      Met Ser Ala Ala Lys Ile Ala Leu Gly Ser Arg Asp Met Cys Ala His
40
                                       25
                                                           30
                  20
      Leu Leu Ile Pro Leu Asn Lys Cys Arg Gln Ala Glu Phe Tyr Leu Pro
                                   40
      Trp Lys Cys Glu Asp Glu Arg His Val Tyr Glu Lys Cys Glu Tyr Glu
45
          50
                              55
                                                   60
      Leu Val Met Glu Arg Met Leu Ala Met Lys Lys Ile Arg Glu Glu Glu
                          7.0
                                              75
      Ala Leu Ala Lys Gln Asn Lys Leu Gln Gly Asn Ala Ala Val Pro Leu
                      85
                                           90
50
      Ile Pro Lys Thr Ala Asn Ala
                  100
      (2) INFORMATION FOR SEQ ID NO:252:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 94 amino acids
55
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
60
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..94
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1448187
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:252:
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                                     25
      Gln Ala Glu Phe Tyr Leu Pro Trp Lys Cys Glu Asp Glu Arg His Val
      Tyr Glu Lys Cys Glu Tyr Glu Leu Val Met Glu Arg Met Leu Ala Met
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10
      Lys Lys Ile Arg Glu Glu Glu Ala Leu Ala Lys Gln Asn Lys Leu Gln
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                          70
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      (2) INFORMATION FOR SEQ ID NO:253:
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                (A) LENGTH: 87 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
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          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..87
                (D) OTHER INFORMATION: / Ceres Seq. ID 1448188
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          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:253:
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      Leu Leu Ile Pro Leu Asn Lys Cys Arg Gln Ala Glu Phe Tyr Leu Pro
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                                      25
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      Trp Lys Cys Glu Asp Glu Arg His Val Tyr Glu Lys Cys Glu Tyr Glu
                                 40
      Leu Val Met Glu Arg Met Leu Ala Met Lys Lys Ile Arg Glu Glu Glu
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                                                  60
      Ala Leu Ala Lys Gln Asn Lys Leu Gln Gly Asn Ala Ala Val Pro Leu
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      Ile Pro Lys Thr Ala Asn Ala
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      (2) INFORMATION FOR SEQ ID NO:254:
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                (A) LENGTH: 2034 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
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          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..2034
                (D) OTHER INFORMATION: / Ceres Seq. ID 1450875
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:254:
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                                                                             180
      aaactctagt aagtctaaag cagagcctga tgagcctaaa ggatcgggtt ttgatcctga
                                                                             240
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                                                                             300
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                                                                             360
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                                                                             420
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                                                                             480
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                                                                             600
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      aqaaqaacaq atccaaqctc aqcatcqcca aactqaqaaa qaqaqaqctq aacttqaqcq
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      agagacgatt cgtgtcaagg ccatggctga agctgaaggt cgggctcatg aagccaaact
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      tactqaaqaq caaaacaqaa qattgcttat ggaaaggatt aatggtgaaa gagagaagtg
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_																caatt		
5																gcttga		
																acagc ggtac		
						_				_	_					.gg.ac :atgat		
							_	_	_	_	_	_			_	gatatt		
10	_		-	_	-			_		_	_				_	gcttt	1380	
																gcttt		
			_	-	_				_				_		_	agacc		
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		_														gacaga		
	aggattetet ggeegtgaga ttgeaaaget tgtggetgga gtacaagetg gagtgtaegg aegageggat tgtgttttgg atteaeaget ttttaaagag attgttgaat ataaggttga																	
	_				_	-		_				_	_		-			
20																ctctta		
20	2 22 2						acatgicatt tactigatea gaattittag attaaacatg						gaaagaagaa totgattgtt					
		INFC	-					_		aaac	acy	Llac	ayay	gic i	Llac			
	( 2 )		SEÇ															
		( 1 )					39 am			ic								
25							no ac		2010	10								
					RANI													
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		(ii)	MOI															
		(ix)	FEP	TURE	: :													
30							pept											
							16											
	(D) OTHER INFORMATION: / Ceres Seq. ID 1450876																	
										ID MC								
2.5		Ser	Lys	Ser		Ala	Ala	Ser	Arg	Leu	Cys	Ser	Ala	Ala	Ala	Ile		
35	1									7.0					4.5			
				D.1	5	<u> </u>		<b>a</b> -		10	Q1	70	70	<b>7</b> . 7	15	7.7		
	Ala	Ala	Ala		-	Ser	Met	Ser			G1n	Asn	Arg			Ala		
				20	Thr				25	Ser			_	30	Tyr			
		Ala Ser	Ser	20	Thr			Pro	25	Ser			Ser	30	Tyr			
4 N	Asp	Ser	Ser 35	20 Arg	Thr	Arg	Phe	Pro 40	25 Phe	Ser Phe	Ser	Ser	Ser 45	30 Pro	Tyr Ser	Pro		
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40 45	Asp Pro Ser 65	Ser Pro 50 Lys	Ser 35 Ser Ala	20 Arg Asp Glu	Thr Phe Ser Pro	Arg Pro Asp 70	Phe Ala 55 Glu	Pro 40 Asn Pro	25 Phe Gln Lys	Ser Phe Ser Gly	Ser Ser Ser 75	Ser Thr 60 Gly	Ser 45 Asn Phe	30 Pro Ser Asp	Tyr Ser Ser Pro	Pro Lys Glu 80		
	Asp Pro Ser 65 Ala	Ser Pro 50 Lys	Ser 35 Ser Ala Glu	20 Arg Asp Glu Arg	Thr Phe Ser Pro Ala 85	Arg Pro Asp 70 Ala	Phe Ala 55 Glu Lys	Pro 40 Asn Pro Ala	25 Phe Gln Lys Leu	Ser Phe Ser Gly Arg 90	Ser Ser Ser 75 Asp	Ser Thr 60 Gly Ile	Ser 45 Asn Phe Asn	30 Pro Ser Asp	Tyr Ser Ser Pro Ser 95	Pro Lys Glu 80 Pro		
	Asp Pro Ser 65 Ala	Ser Pro 50 Lys Leu	Ser 35 Ser Ala Glu	20 Arg Asp Glu Arg	Thr Phe Ser Pro Ala 85	Arg Pro Asp 70 Ala	Phe Ala 55 Glu Lys	Pro 40 Asn Pro Ala	25 Phe Gln Lys Leu	Ser Phe Ser Gly Arg 90	Ser Ser Ser 75 Asp	Ser Thr 60 Gly Ile	Ser 45 Asn Phe Asn	30 Pro Ser Asp	Tyr Ser Ser Pro Ser 95	Pro Lys Glu 80 Pro		
	Asp Pro Ser 65 Ala His	Ser Pro 50 Lys Leu	Ser 35 Ser Ala Glu Lys Glu	20 Arg Asp Glu Arg Gln 100	Thr Phe Ser Pro Ala 85 Val	Arg Pro Asp 70 Ala Phe	Phe Ala 55 Glu Lys Asp	Pro 40 Asn Pro Ala Leu Thr	25 Phe Gln Lys Leu Met 105	Ser Phe Ser Gly Arg 90 Arg	Ser Ser Ser 75 Asp	Ser Thr 60 Gly Ile Gln	Ser 45 Asn Phe Asn Glu Ala	30 Pro Ser Asp Ser Lys 110	Tyr Ser Ser Pro Ser 95 Thr	Pro Lys Glu 80 Pro Arg		
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	Asp Pro Ser 65 Ala His Leu	Ser Pro 50 Lys Leu Ser Ala Asn	Ser 35 Ser Ala Glu Lys Glu 115	20 Arg Asp Glu Arg Gln 100 Leu	Thr Phe Ser Pro Ala 85 Val Ala	Arg Pro Asp 70 Ala Phe Ala	Phe Ala 55 Glu Lys Asp Glu Gln	Pro 40 Asn Pro Ala Leu Thr 120	25 Phe Gln Lys Leu Met 105 Ser	Ser Phe Ser Gly Arg 90 Arg	Ser Ser 75 Asp Lys Tyr	Ser Thr 60 Gly Ile Gln Glu Glu	Ser 45 Asn Phe Asn Glu Ala 125	30 Pro Ser Asp Ser Lys 110 Ile	Ser Ser Pro Ser 95 Thr	Pro Lys Glu 80 Pro Arg		
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45	Asp Pro Ser 65 Ala His Leu His	Ser Pro 50 Lys Leu Ser Ala Asn	Ser 35 Ser Ala Glu Lys Glu 115 Asp	20 Arg Asp Glu Arg Gln 100 Leu Ile	Thr Phe Ser Pro Ala 85 Val Ala Gly	Arg Pro Asp 70 Ala Phe Ala Arg Ala	Phe Ala 55 Glu Lys Asp Glu Gln 135	Pro 40 Asn Pro Ala Leu Thr 120 Gln	25 Phe Gln Lys Leu Met 105 Ser Lys	Ser Phe Ser Gly Arg 90 Arg His	Ser Ser 75 Asp Lys Tyr Ala Gln	Ser Thr 60 Gly Ile Gln Glu Glu 140	Ser 45 Asn Phe Asn Glu Ala 125 Asp	30 Pro Ser Asp Ser Lys 110 Ile	Tyr Ser Ser Pro Ser 95 Thr Gln Arg	Pro Lys Glu 80 Pro Arg Ala Asn Glu		
45	Asp Pro Ser 65 Ala His Leu His	Ser Pro 50 Lys Leu Ser Ala Asn 130 Leu	Ser 35 Ser Ala Glu Lys Glu 115 Asp	20 Arg Asp Glu Arg Gln 100 Leu Ile Thr	Thr Phe Ser Pro Ala 85 Val Ala Gly Gln	Arg Pro Asp 70 Ala Phe Ala Arg Ala 150	Phe Ala 55 Glu Lys Asp Glu Gln 135 Gln	Pro 40 Asn Pro Ala Leu Thr 120 Gln	25 Phe Gln Lys Leu Met 105 Ser Lys	Ser Phe Ser Gly Arg 90 Arg His Leu	Ser Ser 75 Asp Lys Tyr Ala Gln 155	Ser Thr 60 Gly Ile Gln Glu Glu 140 Asn	Ser 45 Asn Phe Asn Glu Ala 125 Asp Leu	30 Pro Ser Asp Ser Lys 110 Ile Gln Arg	Tyr Ser Ser Pro Ser 95 Thr Gln Arg	Pro Lys Glu 80 Pro Arg Ala Asn Glu 160		
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45 50	Asp Pro Ser 65 Ala His Leu 145 Asp	Ser Pro 50 Lys Leu Ser Ala Asn 130 Leu Glu His	Ser 35 Ser Ala Glu Lys Glu 115 Asp Gln Leu Asn	20 Arg Asp Glu Arg Gln 100 Leu Ile Thr Ala Val 180	Thr Phe Ser Pro Ala 85 Val Ala Gly Gln Arg 165 Glu	Arg Pro Asp 70 Ala Phe Ala Arg Ala 150 Lys Leu	Phe Ala 55 Glu Lys Asp Glu Gln 135 Gln Arg Val	Pro 40 Asn Pro Ala Leu Thr 120 Gln Thr Gln Lys	25 Phe Gln Lys Leu Met 105 Ser Lys Gln Met 185	Ser Phe Ser Gly Arg 90 Arg His Leu Ala Thr 170 Gln	Ser Ser 75 Asp Lys Tyr Ala Gln 155 Asp Glu	Ser Thr 60 Gly Ile Gln Glu 140 Asn His Ala	Ser 45 Asn Phe Asn Glu Ala 125 Asp Leu Glu Ser	30 Pro Ser Asp Ser Lys 110 Ile Gln Arg Ala Ser 190	Ser Ser Pro Ser 95 Thr Gln Arg Tyr Gln 175 Ile	Pro Lys Glu 80 Pro Arg Ala Asn Glu 160 Arg Arg		
45 50	Asp Pro Ser 65 Ala His Leu 145 Asp	Ser Pro 50 Lys Leu Ser Ala Asn 130 Leu Glu	Ser 35 Ser Ala Glu Lys Glu 115 Asp Gln Leu Asn	20 Arg Asp Glu Arg Gln 100 Leu Ile Thr Ala Val 180	Thr Phe Ser Pro Ala 85 Val Ala Gly Gln Arg 165 Glu	Arg Pro Asp 70 Ala Phe Ala Arg Ala 150 Lys Leu	Phe Ala 55 Glu Lys Asp Glu Gln 135 Gln Arg Val	Pro 40 Asn Pro Ala Leu Thr 120 Gln Thr Gln Lys	25 Phe Gln Lys Leu Met 105 Ser Lys Gln Met 185	Ser Phe Ser Gly Arg 90 Arg His Leu Ala Thr 170 Gln	Ser Ser 75 Asp Lys Tyr Ala Gln 155 Asp Glu	Ser Thr 60 Gly Ile Gln Glu 140 Asn His Ala	Ser 45 Asn Phe Asn Glu Ala 125 Asp Leu Glu Ser	30 Pro Ser Asp Ser Lys 110 Ile Gln Arg Ala Ser 190	Ser Ser Pro Ser 95 Thr Gln Arg Tyr Gln 175 Ile	Pro Lys Glu 80 Pro Arg Ala Asn Glu 160 Arg Arg		
45 50 55	Asp Pro Ser 65 Ala His Leu His Leu 145 Asp	Ser Pro 50 Lys Leu Ser Ala Asn 130 Leu Glu His Glu	Ser 35 Ser Ala Glu Lys Glu 115 Asp Gln Leu Asn	20 Arg Asp Glu Arg Gln 100 Leu Ile Thr Ala Val 180 Ala	Thr Phe Ser Pro Ala 85 Val Ala Gly Gln Arg 165 Glu Lys	Arg Pro Asp 70 Ala Phe Ala Arg Ala 150 Lys Leu Ile	Phe Ala 55 Glu Lys Asp Glu Gln 135 Gln Arg Val Ala	Pro 40 Asn Pro Ala Leu Thr 120 Gln Lys Thr 200	25 Phe Gln Lys Leu Met 105 Ser Lys Gln Met 185 Glu	Ser Phe Ser Gly Arg 90 Arg His Leu Ala Thr 170 Gln	Ser Ser 75 Asp Lys Tyr Ala Gln 155 Asp Glu Gln	Ser Thr 60 Gly Ile Gln Glu 140 Asn His Ala Ile	Ser 45 Asn Phe Asn Glu Ala 125 Asp Leu Glu Ser Gln 205	Ser Asp Ser Lys 110 Ile Gln Arg Ala Ser 190 Ala	Ser Ser Pro Ser 95 Thr Gln Arg Tyr Gln 175 Ile Gln	Pro Lys Glu 80 Pro Arg Ala Asn Glu 160 Arg Arg His		
45 50	Asp Pro Ser 65 Ala His Leu His Leu 145 Asp	Ser Pro 50 Lys Leu Ser Ala Asn 130 Leu Glu His	Ser 35 Ser Ala Glu Lys Glu 115 Asp Gln Leu Asn	20 Arg Asp Glu Arg Gln 100 Leu Ile Thr Ala Val 180 Ala	Thr Phe Ser Pro Ala 85 Val Ala Gly Gln Arg 165 Glu Lys	Arg Pro Asp 70 Ala Phe Ala Arg Ala 150 Lys Leu Ile	Phe Ala 55 Glu Lys Asp Glu Gln 135 Gln Arg Val Ala	Pro 40 Asn Pro Ala Leu Thr 120 Gln Lys Thr 200	25 Phe Gln Lys Leu Met 105 Ser Lys Gln Met 185 Glu	Ser Phe Ser Gly Arg 90 Arg His Leu Ala Thr 170 Gln	Ser Ser 75 Asp Lys Tyr Ala Gln 155 Asp Glu Gln	Ser Thr 60 Gly Ile Gln Glu 140 Asn His Ala Ile	Ser 45 Asn Phe Asn Glu Ala 125 Asp Leu Glu Ser Gln 205 Glu	Ser Asp Ser Lys 110 Ile Gln Arg Ala Ser 190 Ala	Ser Ser Pro Ser 95 Thr Gln Arg Tyr Gln 175 Ile Gln	Pro Lys Glu 80 Pro Arg Ala Asn Glu 160 Arg Arg His		

Val Lys Ala Met Ala Glu Ala Glu Gly Arg Ala His Glu Ala Lys Leu

443 230 235 Thr Glu Glu Gln Asn Arg Arg Leu Leu Met Glu Arg Ile Asn Gly Glu 250 255 245 Arg Glu Lys Trp Leu Ala Ala Ile Asn Thr Met Phe Ser His Ile Glu 5 260 265 270 Gly Gly Phe Arg Thr Leu Leu Thr Asp Arg Asn Lys Leu Ile Met Thr 280 285 Val Gly Gly Ala Thr Ala Leu Ala Ala Gly Val Tyr Thr Thr Arg Glu 295 300 10 Gly Ala Arg Val Thr Trp Gly Tyr Ile Asn Arg Met Leu Gly Gln Pro 310 315 320 Ser Leu Ile Arg Glu Ser Ser Met Arg Arg Phe Pro Trp Thr Gly Ser Val Ser Gln Phe Lys Asn Arg Ile Ser Gly Ala Ala Ala Ser Ala 15 345 Ala Glu Gly Lys Lys Pro Leu Asp Asn Val Ile Leu His Thr Ser Leu 360 Lys Lys Arg Ile Glu Arg Leu Ala Arg Ala Thr Ala Asn Thr Lys Ser 375 20 His Gln Ala Pro Phe Arg Asn Met Met Phe Tyr Gly Pro Pro Gly Thr 390 395 400 Gly Lys Thr Met Val Ala Arg Glu Ile Ala Arg Lys Ser Gly Leu Asp 410 405 Tyr Ala Met Met Thr Gly Gly Asp Val Ala Pro Leu Gly Ser Gln Ala 25 420 425 Val Thr Lys Ile His Gln Ile Phe Asp Trp Ala Lys Lys Ser Asn Arg 440 Gly Leu Leu Phe Ile Asp Glu Ala Asp Ala Phe Leu Cys Glu Arg 455 460 30 Asn Ser Thr Tyr Met Ser Glu Ala Gln Arg Ser Ala Leu Asn Ala Leu 470 475 Leu Phe Arg Thr Gly Asp Gln Ser Arg Asp Ile Val Leu Val Leu Ala 485 490 495 Thr Asn Arg Pro Gly Asp Leu Asp Ser Ala Val Thr Asp Arg Ile Asp 35 500 505 510 Glu Val Ile Glu Phe Pro Leu Pro Gly Glu Glu Glu Arg Phe Lys Leu 520 Leu Asn Leu Tyr Leu Asn Lys Tyr Leu Lys Met Gly Asp Asn Asn Glu 535 40 Asp Thr Lys Pro Lys Trp Ser His Leu Phe Lys Lys Leu Ser Gln Lys 550 555 560 Ile Thr Val Glu Glu Asp Leu Thr Asp Lys Val Ile Ser Glu Ala Ala 570 575 565 Lys Lys Thr Glu Gly Phe Ser Gly Arg Glu Ile Ala Lys Leu Val Ala 45 585 590 Gly Val Gln Ala Gly Val Tyr Gly Arg Ala Asp Cys Val Leu Asp Ser 600 Gln Leu Phe Lys Glu Ile Val Glu Tyr Lys Val Glu Glu His His Arg 610 615 620 50 Arg His Met Leu Ala Ser Glu Gly Phe Gln Pro Leu Leu Phe Ser 630 635 (2) INFORMATION FOR SEQ ID NO:256: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 635 amino acids 55 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 60 (A) NAME/KEY: peptide (B) LOCATION: 1..635 (D) OTHER INFORMATION: / Ceres Seq. ID 1450877

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5	Thr	Ser	Met	Ser 20	Met	Ser	Gln	Asn	Arg 25	Ala	Tyr	Ala	Asp	Ser 30	Ser	Arg
	Phe	Arg	Phe 35	Pro	Phe	Phe	Ser	Ser 40	Ser	Pro	Ser	Pro	Pro 45	Pro	Ser	Asp
10		50			Gln		55					60		_		
	65	_			Lys	70					75					80
			_		Leu 85					90					95	
15			_	100	Met	_			105					110		
			115		Ser			120					125			
20 25 30	_	130			Lys		135		_		_	140				
	145				Lys	150					155					160
	-	-	_		Gln 165		_			170		_			175	
				180	Met				185					190	_	
	_		195		Glu Glu			200					205			
	_	210	_		Gly		215					220		_		
	225				Leu	230					235					240
35		-	_		245 Asn					250	_				255	_
				260	Asp				265				_	270		_
			275		Ala			280					285	_	-	
		290				_	295					300			-	Arg
	305				Arg	310					315					320
45 50					325 Ser	-			_	330	_				335	
	Lys	Pro	Leu	340 Asp	Asn	Val	Ile	Leu	345 His	Thr	Ser	Leu	Lys	350 Lys	Arg	Ile
	Glu	Arg	355 Leu	Ala	Arg	Ala	Thr	360 Ala	Asn	Thr	Lys	Ser	365 His	Gln	Ala	Pro
	Phe	370 Arg	Asn	Met	Met	Phe	375 Tyr	Gly	Pro	Pro	Gly	380 Thr	Gly	Lys	Thr	Met
	385 Val	Ala	Arg	Glu	Ile	390 Ala	Arg	Lys	Ser	Gly	395 Leu	Asp	Tyr	Ala	Met	400 Met
55	Thr	Gly	Gly		405 Val	Ala	Pro	Leu		410 Ser	Gln	Ala	Val		415 Lys	Ile
	His	Gln		420 Phe	Asp	Trp	Ala		425 Lys	Ser	Asn	Arg		430 Leu	Leu	Leu
	Phe		435 Asp	Glu	Ala	Asp		440 Phe	Leu	Cys	Glu		445 Asn	Ser	Thr	Tyr
60	Met 465	450 Ser	Glu	Ala	Gln		455 Ser	Ala	Leu	Asn	Ala 475	460 Leu	Leu	Phe	Arg	
		Asp	Gln	Ser	Arg	470 Asp	Ile	Val	Leu	Val		Ala	Thr	Asn	Arg	480 Pro

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     Phe Pro Leu Pro Gly Glu Glu Glu Arg Phe Lys Leu Leu Asn Leu Tyr
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     Lys Trp Ser His Leu Phe Lys Lys Leu Ser Gln Lys Ile Thr Val Glu
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                                           555
     Glu Asp Leu Thr Asp Lys Val Ile Ser Glu Ala Ala Lys Lys Thr Glu
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                    565
     Gly Phe Ser Gly Arg Glu Ile Ala Lys Leu Val Ala Gly Val Gln Ala
                      585
     Gly Val Tyr Gly Arg Ala Asp Cys Val Leu Asp Ser Gln Leu Phe Lys
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                              600
     Glu Ile Val Glu Tyr Lys Val Glu Glu His His Arg Arg His Met Leu
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     Ala Ser Glu Gly Phe Gln Pro Leu Leu Phe Ser
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     (2) INFORMATION FOR SEQ ID NO:257:
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               (A) LENGTH: 617 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
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               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..617
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               (D) OTHER INFORMATION: / Ceres Seq. ID 1450878
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     Ala Asn Gln Ser Ser Thr Asn Ser Ser Lys Ser Lys Ala Glu Pro Asp
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     Glu Pro Lys Gly Ser Gly Phe Asp Pro Glu Ala Leu Glu Arg Ala Ala
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     Lys Ala Leu Arg Asp Ile Asn Ser Ser Pro His Ser Lys Gln Val Phe
                                           75
     Asp Leu Met Arg Lys Gln Glu Lys Thr Arg Leu Ala Glu Leu Ala Ala
      Glu Thr Ser His Tyr Glu Ala Ile Gln Ala His Asn Asp Ile Gly Arg
                                   105
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      Gln Thr Lys Ala Gln Asn Leu Arg Tyr Glu Asp Glu Leu Ala Arg Lys
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      Val Lys Met Gln Glu Ala Ser Ser Ile Arg Lys Glu Lys Ala Lys Ile
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      Ala Thr Glu Glu Gln Ile Gln Ala Gln His Arg Gln Thr Glu Lys Glu
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      Ala Glu Gly Arg Ala His Glu Ala Lys Leu Thr Glu Glu Gln Asn Arg
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                        230
                                           235
      Ala Ile Asn Thr Met Phe Ser His Ile Glu Gly Gly Phe Arg Thr Leu
```

```
446
                     245
                                        250
                                                            255
     Leu Thr Asp Arg Asn Lys Leu Ile Met Thr Val Gly Gly Ala Thr Ala
                                     265
     Leu Ala Ala Gly Val Tyr Thr Thr Arg Glu Gly Ala Arg Val Thr Trp
 5
                                 280
     Gly Tyr Ile Asn Arg Met Leu Gly Gln Pro Ser Leu Ile Arg Glu Ser
                         295
     Ser Met Arg Arg Phe Pro Trp Thr Gly Ser Val Ser Gln Phe Lys Asn
                         310
                                            315
10
     Arg Ile Ser Gly Ala Ala Ala Ala Ser Ala Ala Glu Gly Lys Lys Pro
                                        330
     Leu Asp Asn Val Ile Leu His Thr Ser Leu Lys Lys Arg Ile Glu Arg
                                    345
     Leu Ala Arg Ala Thr Ala Asn Thr Lys Ser His Gln Ala Pro Phe Arg
15
                                 360
     Asn Met Met Phe Tyr Gly Pro Pro Gly Thr Gly Lys Thr Met Val Ala
                             375
     Arg Glu Ile Ala Arg Lys Ser Gly Leu Asp Tyr Ala Met Met Thr Gly
                         390
                                            395
20
     Gly Asp Val Ala Pro Leu Gly Ser Gln Ala Val Thr Lys Ile His Gln
                                        410
                     405
     Ile Phe Asp Trp Ala Lys Lys Ser Asn Arg Gly Leu Leu Leu Phe Ile
                                    425
                 420
     Asp Glu Ala Asp Ala Phe Leu Cys Glu Arg Asn Ser Thr Tyr Met Ser
25
                             440
                                                    445
     Glu Ala Gln Arg Ser Ala Leu Asn Ala Leu Leu Phe Arg Thr Gly Asp
                             455
                                                460
     Gln Ser Arg Asp Ile Val Leu Val Leu Ala Thr Asn Arg Pro Gly Asp
                         470
                                            475
30
     Leu Asp Ser Ala Val Thr Asp Arg Ile Asp Glu Val Ile Glu Phe Pro
                                        490
     Leu Pro Gly Glu Glu Arg Phe Lys Leu Leu Asn Leu Tyr Leu Asn
                                    505
     Lys Tyr Leu Lys Met Gly Asp Asn Asn Glu Asp Thr Lys Pro Lys Trp
35
                                 520
      Ser His Leu Phe Lys Lys Leu Ser Gln Lys Ile Thr Val Glu Glu Asp
                             535
                                                540
     Leu Thr Asp Lys Val Ile Ser Glu Ala Ala Lys Lys Thr Glu Gly Phe
                         550
                                            555
     Ser Gly Arg Glu Ile Ala Lys Leu Val Ala Gly Val Gln Ala Gly Val
40
                                        570
                     565
      Tyr Gly Arg Ala Asp Cys Val Leu Asp Ser Gln Leu Phe Lys Glu Ile
                 580
                                    585
      Val Glu Tyr Lys Val Glu Glu His His Arg Arg His Met Leu Ala Ser
45
                                600
             595
      Glu Gly Phe Gln Pro Leu Leu Phe Ser
                            615
          610
      (2) INFORMATION FOR SEQ ID NO:258:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 478 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
55
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..478
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459191
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:258:
60
      aaccctcttg gaaagagtct caacacttgc agagaaaaag aacaaggaag atcccggaaa
      atggcaacgg cgattgtacg ttcagctctt tcccgagcag tgactcgcgc agtccgaaga
      catecqteqe tectaaqeqa aaetttteet etteegeegg eeatgaegat gettatgaag
```

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ctgcgaagtg ggagaagata acttatctgg gtattgctag ttgcactgct ctagctgtct
      atgttttatc caagggccat catcacggcg aagacaagga gtttccttgg ggtccggatg
                                                                             300
      gtctgtttga ggtgaagcac aacaaagagc actgagtctt gcgtggtcat aataacgtct
                                                                             360
     tctagqttta tttgaaaqgc taaaatgttt taccgtattt gttctcaccg tttgtcaacg
                                                                             420
 5
      atttgctact ccaatctctt ttcttttgtt gggaaataaa agttaatact ttgcttgg
      (2) INFORMATION FOR SEQ ID NO:259:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 66 amino acids
                (B) TYPE: amino acid
10
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
15
                (B) LOCATION: 1..66
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459192
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:259:
      Asn Pro Leu Gly Lys Ser Leu Asn Thr Cys Arg Glu Lys Glu Gln Gly
                                          10
20
     Arg Ser Arg Lys Met Ala Thr Ala Ile Val Arg Ser Ala Leu Ser Arg
      Ala Val Thr Arg Ala Val Arg Arg His Pro Ser Leu Leu Ser Glu Thr
                                  40
      Phe Pro Leu Pro Pro Ala Met Thr Met Leu Met Lys Leu Arg Ser Gly
25
      Arg Arg
      65
      (2) INFORMATION FOR SEQ ID NO:260:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 110 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..110
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459193
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:260:
40
      Pro Ser Trp Lys Glu Ser Gln His Leu Gln Arg Lys Arg Thr Arg Lys
                                          10
      Ile Pro Glu Asn Gly Asn Gly Asp Cys Thr Phe Ser Ser Phe Pro Ser
      Ser Asp Ser Arg Ser Pro Lys Thr Ser Val Ala Pro Lys Arg Asn Phe
45
                                  40
      Ser Ser Ser Ala Gly His Asp Asp Ala Tyr Glu Ala Ala Lys Trp Glu
      Lys Ile Thr Tyr Leu Gly Ile Ala Ser Cys Thr Ala Leu Ala Val Tyr
                          70
                                               75
      Val Leu Ser Lys Gly His His His Gly Glu Asp Lys Glu Phe Pro Trp
50
                                          90
      Gly Pro Asp Gly Leu Phe Glu Val Lys His Asn Lys Glu His
                                      105
                  100
      (2) INFORMATION FOR SEQ ID NO:261:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 741 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
```

(A) NAME/KEY: -

```
(B) LOCATION: 1..741
                (D) OTHER INFORMATION: / Ceres Seq. ID 1461848
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:261:
     attttagtac attgttgacc atctttttcg tatagactac tatctctgat ctcttgcgag
                                                                             60
5
     ttaaqtcagt aactaggaaa attcagaagc gctctcaatc tcaaaaatat ccatggcggc
                                                                            120
     gattacagaa tttctaccaa aagagtacgg atatgtcgtt ctcgtcctcg tcttctactg
                                                                            180
     tttcctcaac ctctggatgg gtgctcaagt cggcagagct cgcaaaaaggt acaacgtccc
                                                                            240
     gtatccaact ctatatgcaa tagaatcaga aaacaaagat gctaagctct tcaactgtgt
                                                                            300
     tcagagagga catcaaaact ctttagagat gatgccaatg tatttcatac tgatgatcct
                                                                            360
10
     cggtgggatg aagcaccett gtatetgtae tggeettggt ttgetttaea aegttageeg
                                                                            420
     attettetae tttaaaggtt atgetaetgg agateeeatg aagegtetta egategggaa
                                                                            480
     atacggtttc ttggggttgc taggtctgat gatatgtacc atctcgtttg gtgtcactct
                                                                            540
     gatccttgct taagctactc gtttctgggg ttaatgattc tctggtttgc tcgaagaata
                                                                            600
     tagaaccaat gcttgtaagc tgtccacaaa acttgtgtaa tactttagag tttgtcactt
                                                                            660
15
     ttaaaagttt gtaataaatc atggcttcat agaacagttg aaatttcaca tccgtagacg
                                                                            720
     ttaataaaga tttgaattat q
     (2) INFORMATION FOR SEQ ID NO:262:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 146 amino acids
20
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
25
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..146
                (D) OTHER INFORMATION: / Ceres Seq. ID 1461849
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:262:
     Met Ala Ala Ile Thr Glu Phe Leu Pro Lys Glu Tyr Gly Tyr Val Val
30
                                         10
     Leu Val Leu Val Phe Tyr Cys Phe Leu Asn Leu Trp Met Gly Ala Gln
                                     25
     Val Gly Arg Ala Arg Lys Arg Tyr Asn Val Pro Tyr Pro Thr Leu Tyr
                                  40
35
     Ala Ile Glu Ser Glu Asn Lys Asp Ala Lys Leu Phe Asn Cys Val Gln
                              5.5
     Arg Gly His Gln Asn Ser Leu Glu Met Met Pro Met Tyr Phe Ile Leu
                                              75
                          70
     Met Ile Leu Gly Gly Met Lys His Pro Cys Ile Cys Thr Gly Leu Gly
40
                      8.5
                                          90
      Leu Leu Tyr Asn Val Ser Arg Phe Phe Tyr Phe Lys Gly Tyr Ala Thr
                                     105
                  100
                                                          110
      Gly Asp Pro Met Lys Arg Leu Thr Ile Gly Lys Tyr Gly Phe Leu Gly
                              120
                                                      125
45
      Leu Leu Gly Leu Met Ile Cys Thr Ile Ser Phe Gly Val Thr Leu Ile
                              135
          130
                                                 140
      Leu Ala
      145
      (2) INFORMATION FOR SEQ ID NO:263:
50
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 118 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
55
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..118
                (D) OTHER INFORMATION: / Ceres Seq. ID 1461850
60
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:263:
      Met Gly Ala Gln Val Gly Arg Ala Arg Lys Arg Tyr Asn Val Pro Tyr
                                          10
```

449 Pro Thr Leu Tyr Ala Ile Glu Ser Glu Asn Lys Asp Ala Lys Leu Phe 25 Asn Cys Val Gln Arg Gly His Gln Asn Ser Leu Glu Met Met Pro Met 40 5 Tyr Phe Ile Leu Met Ile Leu Gly Gly Met Lys His Pro Cys Ile Cys 55 Thr Gly Leu Gly Leu Leu Tyr Asn Val Ser Arg Phe Phe Tyr Phe Lys 70 75 Gly Tyr Ala Thr Gly Asp Pro Met Lys Arg Leu Thr Ile Gly Lys Tyr 10 90 Gly Phe Leu Gly Leu Gly Leu Met Ile Cys Thr Ile Ser Phe Gly 105 100 Val Thr Leu Ile Leu Ala 115 15 (2) INFORMATION FOR SEQ ID NO:264: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 74 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..74 25 (D) OTHER INFORMATION: / Ceres Seq. ID 1461851 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:264: Met Met Pro Met Tyr Phe Ile Leu Met Ile Leu Gly Gly Met Lys His 10 Pro Cys Ile Cys Thr Gly Leu Gly Leu Leu Tyr Asn Val Ser Arg Phe 30 20 25 Phe Tyr Phe Lys Gly Tyr Ala Thr Gly Asp Pro Met Lys Arg Leu Thr 40 Ile Gly Lys Tyr Gly Phe Leu Gly Leu Gly Leu Met Ile Cys Thr 55 35 Ile Ser Phe Gly Val Thr Leu Ile Leu Ala 7.0 (2) INFORMATION FOR SEQ ID NO: 265: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 469 base pairs 40 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 45 (A) NAME/KEY: -(B) LOCATION: 1..469 (D) OTHER INFORMATION: / Ceres Seq. ID 1472772 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:265: atataqaata taaccatatt ccggacatga agatcgtgac attggtactc gtcgtcttcg 50 tcatactttc gacatcattc ccggctgcca tcaaagccga agacacggga gatacaggaa 120 atgtgggagt gacatgtgac gcaaggcagc ttcagccttg ccttgccgcg attacgggag 180 gaggacaacc ctcgggtgca tgttgtgcaa agcttacaga gcaacagtca tgcctatgtg 240 qtttcqctaa qaaccctqcq ttcqcacaqt acattaqctc tccaaacqct cqcaaaqtqc 300 tccttgcttg caatgttgct tatcccactt gttgaaactt atctagattt tataaataaa 360 55 taaacqaaaq gaaataaatt acattatatc aaacgttatg atacaattca accgtttgtg 420 ttaatgtact ggcttacatg gttaaataaa gtttaatttc ttggttgcc (2) INFORMATION FOR SEQ ID NO: 266: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 110 amino acids

> (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear

```
(ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..110
 5
                (D) OTHER INFORMATION: / Ceres Seq. ID 1472773
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:266:
     Ile Glu Tyr Asn His Ile Pro Asp Met Lys Ile Val Thr Leu Val Leu
                                          10
     Val Val Phe Val Ile Leu Ser Thr Ser Phe Pro Ala Ala Ile Lys Ala
10
                  20
                                      25
     Glu Asp Thr Gly Asp Thr Gly Asn Val Gly Val Thr Cys Asp Ala Arg
                                  40
                                                      45
     Gln Leu Gln Pro Cys Leu Ala Ala Ile Thr Gly Gly Gln Pro Ser
                             55
15
     Gly Ala Cys Cys Ala Lys Leu Thr Glu Gln Gln Ser Cys Leu Cys Gly
                         70
                                              75
      Phe Ala Lys Asn Pro Ala Phe Ala Gln Tyr Ile Ser Ser Pro Asn Ala
                                         90
     Arg Lys Val Leu Leu Ala Cys Asn Val Ala Tyr Pro Thr Cys
20
                 100
      (2) INFORMATION FOR SEQ ID NO:267:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 102 amino acids
                (B) TYPE: amino acid
25
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
30
                (B) LOCATION: 1..102
                (D) OTHER INFORMATION: / Ceres Seq. ID 1472774
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:267:
      Met Lys Ile Val Thr Leu Val Leu Val Val Phe Val Ile Leu Ser Thr
                                          10
35
      Ser Phe Pro Ala Ala Ile Lys Ala Glu Asp Thr Gly Asp Thr Gly Asn
                                      25
      Val Gly Val Thr Cys Asp Ala Arg Gln Leu Gln Pro Cys Leu Ala Ala
                                  40
      Ile Thr Gly Gly Gln Pro Ser Gly Ala Cys Cys Ala Lys Leu Thr
40
      Glu Gln Gln Ser Cys Leu Cys Gly Phe Ala Lys Asn Pro Ala Phe Ala
                          70
                                              75
      Gln Tyr Ile Ser Ser Pro Asn Ala Arg Lys Val Leu Leu Ala Cys Asn
                      8.5
45
      Val Ala Tyr Pro Thr Cys
                  100
      (2) INFORMATION FOR SEQ ID NO:268:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1056 base pairs
50
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
55
                (A) NAME/KEY: -
                (B) LOCATION: 1..1056
                (D) OTHER INFORMATION: / Ceres Seq. ID 1533352
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:268:
      aaaaaaaaaa accaaagcaa aaaaatggct ttgaaactca cttctccgcc ttcagttttc
                                                                              60
60
      tcacaatcaa ggagattatc ttcttcttcg ttaattccga taaggtcaaa atccacattc
                                                                             120
      accqqatttc qatcqaqaac cggtgtttat ttaaqcaaaa cgacggcgct tcagtcgtct
      acaaaactga qtqtqqcqqc ggagagtcct gcggcgacaa ttgcgacgga tgattggqqq
                                                                             240
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aaaqtatcqq cqqttctqtt tqatatqqac qqtqtqcttt qtaacaqtqa agatctttct
                                                                          300
     agacgcgccg ccgtggatgt ttttacggag atgggagttg aagtcactgt ggacgatttc
                                                                          360
     gttcctttta tgggaacagg tgaagccaag tttttaggag gtgttgcttc agtcaaagaa
                                                                          420
     gttaaaggat ttgatccaga tgcagctaaa aagagattct ttgaaatata tctcgataag
                                                                          480
 5
     tatgcgaagc cagaatctgg gattggattt ccaggagcat tggagcttgt tactgagtgt
                                                                          540
     aagaacaaag gccttaaagt cgctgttgca tctagtgctg accgtatcaa agttgatgcg
                                                                          600
     aatctgaaag ctgctggttt gtctttgacc atgtttgatg ccattgtttc agcagatgcc
                                                                          660
     tttgagaatt tgaaaccagc tccagatatt ttcctggctg ctgcaaagat cttaggtgtg
                                                                          720
     cctaccagcg agtgtgttgt tattgaagat gcgcttgctg gagtccaagc cgcacaagct
                                                                          780
10
     gcgaacatga gatgtatagc cgtaaaaact actttatctg aagcaattct taaggatgct
                                                                          840
     ggtccttcaa tgatacgaga cgatattgga aacatctcaa tcaatgacat tctcactggt
                                                                          900
     ggctcagatt ctaccagtat gtagtctcaa agaaattcga tggaaaatat cgttcttttc
                                                                         960
     atgtgtattt tatttcttgt ttactccttt tgaaaacttt tgaataaagg ggctttcttt
                                                                         1020
     gtaacgagat tacacattta aaacaatctt ttctgt
15
     (2) INFORMATION FOR SEQ ID NO: 269:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 307 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
20
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..307
25
               (D) OTHER INFORMATION: / Ceres Seq. ID 1533353
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:269:
     Lys Lys Lys Asn Gln Ser Lys Lys Met Ala Leu Lys Leu Thr Ser Pro
                                        1.0
     Pro Ser Val Phe Ser Gln Ser Arg Arg Leu Ser Ser Ser Leu Ile
30
                 2.0
                                     25
     Pro Ile Arg Ser Lys Ser Thr Phe Thr Gly Phe Arg Ser Arg Thr Gly
                                 40
     Val Tyr Leu Ser Lys Thr Thr Ala Leu Gln Ser Ser Thr Lys Leu Ser
                             55
35
     Val Ala Ala Glu Ser Pro Ala Ala Thr Ile Ala Thr Asp Asp Trp Gly
                         70
                                             75
     Lys Val Ser Ala Val Leu Phe Asp Met Asp Gly Val Leu Cys Asn Ser
                                         90
      Glu Asp Leu Ser Arg Arg Ala Ala Val Asp Val Phe Thr Glu Met Gly
40
                                     105
     Val Glu Val Thr Val Asp Asp Phe Val Pro Phe Met Gly Thr Gly Glu
                                 120
                                                    125
     Ala Lys Phe Leu Gly Gly Val Ala Ser Val Lys Glu Val Lys Gly Phe
                             135
                                               140
         130
45
     Asp Pro Asp Ala Ala Lys Lys Arg Phe Phe Glu Ile Tyr Leu Asp Lys
                         150
                                            155
      Tyr Ala Lys Pro Glu Ser Gly Ile Gly Phe Pro Gly Ala Leu Glu Leu
                     165
                                         170
      Val Thr Glu Cys Lys Asn Lys Gly Leu Lys Val Ala Val Ala Ser Ser
50
                                     185
                 180
     Ala Asp Arg Ile Lys Val Asp Ala Asn Leu Lys Ala Ala Gly Leu Ser
                                200
                                                    205
      Leu Thr Met Phe Asp Ala Ile Val Ser Ala Asp Ala Phe Glu Asn Leu
                                                220
                             215
55
      Lys Pro Ala Pro Asp Ile Phe Leu Ala Ala Lys Ile Leu Gly Val
                        230
                                           235
      Pro Thr Ser Glu Cys Val Val Ile Glu Asp Ala Leu Ala Gly Val Gln
                                        250
                     245
      Ala Ala Gln Ala Ala Asn Met Arg Cys Ile Ala Val Lys Thr Thr Leu
60
                                     265
      Ser Glu Ala Ile Leu Lys Asp Ala Gly Pro Ser Met Ile Arg Asp Asp
```

280

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Ile Gly Asn Ile Ser Ile Asn Asp Ile Leu Thr Gly Gly Ser Asp Ser
                            295
     Thr Ser Met
     305
 5
     (2) INFORMATION FOR SEQ ID NC:270:
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 299 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
10
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
              (B) LOCATION: 1..299
15
              (D) OTHER INFORMATION: / Ceres Seq. ID 1533354
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:270:
     Met Ala Leu Lys Leu Thr Ser Pro Pro Ser Val Phe Ser Gln Ser Arg
                                      10
     Arg Leu Ser Ser Ser Leu Ile Pro Ile Arg Ser Lys Ser Thr Phe
20
         20
                            25
     Thr Gly Phe Arg Ser Arg Thr Gly Val Tyr Leu Ser Lys Thr Thr Ala
                              40
     Leu Gln Ser Ser Thr Lys Leu Ser Val Ala Ala Glu Ser Pro Ala Ala
                          55
25
     Thr Ile Ala Thr Asp Asp Trp Gly Lys Val Ser Ala Val Leu Phe Asp
                        7.0
                                          75
     Met Asp Gly Val Leu Cys Asn Ser Glu Asp Leu Ser Arg Arg Ala Ala
                                       90
     Val Asp Val Phe Thr Glu Met Gly Val Glu Val Thr Val Asp Asp Phe
30
                                  105
     Val Pro Phe Met Gly Thr Gly Glu Ala Lys Phe Leu Gly Gly Val Ala
                               120
     Ser Val Lys Glu Val Lys Gly Phe Asp Pro Asp Ala Ala Lys Lys Arg
                           135
35
     Phe Phe Glu Ile Tyr Leu Asp Lys Tyr Ala Lys Pro Glu Ser Gly Ile
                        150
                                          155
     Gly Phe Pro Gly Ala Leu Glu Leu Val Thr Glu Cys Lys Asn Lys Gly
                    165
                                       170
     Leu Lys Val Ala Val Ala Ser Ser Ala Asp Arg Ile Lys Val Asp Ala
40
                                   185
     Asn Leu Lys Ala Ala Gly Leu Ser Leu Thr Met Phe Asp Ala Ile Val
                               200
     Ser Ala Asp Ala Phe Glu Asn Leu Lys Pro Ala Pro Asp Ile Phe Leu
                           215
                                              220
45
     Ala Ala Ala Lys Ile Leu Gly Val Pro Thr Ser Glu Cys Val Val Ile
                        230
                                           235
     Glu Asp Ala Leu Ala Gly Val Gln Ala Ala Gln Ala Ala Asn Met Arg
                    245
                                       250
     Cys Ile Ala Val Lys Thr Thr Leu Ser Glu Ala Ile Leu Lys Asp Ala
50
                      265 270
                260
     Gly Pro Ser Met Ile Arg Asp Asp Ile Gly Asn Ile Ser Ile Asn Asp
                            280
                                                   285
      Ile Leu Thr Gly Gly Ser Asp Ser Thr Ser Met
                         295
55
      (2) INFORMATION FOR SEQ ID NO:271:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 219 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
60
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
```

453

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(A) NAME/KEY: peptide
                (B) LOCATION: 1..219
                (D) OTHER INFORMATION: / Ceres Seq. ID 15333355
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:271:
 5
      Met Asp Gly Val Leu Cys Asn Ser Glu Asp Leu Ser Arg Arg Ala Ala
                                          1.0
      Val Asp Val Phe Thr Glu Met Gly Val Glu Val Thr Val Asp Asp Phe
      Val Pro Phe Met Gly Thr Gly Glu Ala Lys Phe Leu Gly Gly Val Ala
10
                                  40
      Ser Val Lys Glu Val Lys Gly Phe Asp Pro Asp Ala Ala Lys Lys Arg
      Phe Phe Glu Ile Tyr Leu Asp Lys Tyr Ala Lys Pro Glu Ser Gly Ile
                                              75
15
      Gly Phe Pro Gly Ala Leu Glu Leu Val Thr Glu Cys Lys Asn Lys Gly
      Leu Lys Val Ala Val Ala Ser Ser Ala Asp Arg Ile Lys Val Asp Ala
                  100
                                      105
                                                          110
      Asn Leu Lys Ala Ala Gly Leu Ser Leu Thr Met Phe Asp Ala Ile Val
20
                                  120
                                                      125
      Ser Ala Asp Ala Phe Glu Asn Leu Lys Pro Ala Pro Asp Ile Phe Leu
                             135
                                                  140
      Ala Ala Ala Lys Ile Leu Gly Val Pro Thr Ser Glu Cys Val Val Ile
                          150
                                              155
25
      Glu Asp Ala Leu Ala Gly Val Gln Ala Ala Gln Ala Ala Asn Met Arg
                      165
                                          170
      Cys Ile Ala Val Lys Thr Thr Leu Ser Glu Ala Ile Leu Lys Asp Ala
                                      185
      Gly Pro Ser Met Ile Arg Asp Asp Ile Gly Asn Ile Ser Ile Asn Asp
30
                                 200
      Ile Leu Thr Gly Gly Ser Asp Ser Thr Ser Met
                             215
      (2) INFORMATION FOR SEQ ID NO:272:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 1420 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
40
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1420
                (D) OTHER INFORMATION: / Ceres Seq. ID 1534544
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:272:
45
      aaagattett ggteteegat eeagteacet gaagattete ggagettett ttgaceacae
                                                                              60
      actttcaatq gcggatgctg tgaacgctca aactccatcg ctctccgagc aatatcattt
                                                                             120
      qqaqaaaqaa qtqaaqcaaq acacaagtqc aaaqcctqtt qaaqtqaaaq aqqtqqcacc
                                                                             180
      agaagttact acacaagctg aagaggttaa gacggagcaa gctaaggaag aatctcctgt
                                                                             240
      tqaqqaaqcq qtttctqtaq ttqaaqaqaa qtctqaatct qctcctqaat caacqqaaqt
                                                                             300
50
      qqcttctqaq qctcctqctq caqcqqaaqa caatqctqaa qaqactcctq ctqctqctqa
                                                                             360
      agaaaataat gacgaaaacg ctagtgaaga agttgctgaa gaaacccctg atgagatcaa
                                                                             420
      gcttgagaca gctcctgctt gatctcatca ggggtttctt ttttttttt ttttttqaat
                                                                             480
      tttttcacca agtgttcgtt cacggcattg cttgtcctcc agtatgagta tctccggtgc
                                                                             540
      tgcagttggt tcaggcagaa atttaagaag agcggtagag tttgggaaaa ctcatgtggt
                                                                             600
55
      taggcctaaa gggaaacatc aagcaactat tgtctggtta catgggcttg gggacaatgg
                                                                             660
      ctcgagctgg tcccagcttt tggagaccct tccccttcca aatatcaaat ggatttgccc
                                                                             720
      gactgctcct tctcaaccaa taagtttatt tggtggtttt ccctccacag cttggtttga
      tgttgtggac atcaatgaag atggacctga tgatatggaa ggattggatg tggctgctgc
                                                                             840
      acatgttgca aatctgttgt cgaatgagcc tgctgacatt aaattaggtg ttggaggatt
                                                                             900
60
      cagcatgggt gcggcgacat ctctatattc tgcaacttgt tttgctctcg gtaaatatgg
                                                                             960
      aaatggcaat ccatacccta tcaatttaag cgcaatcata ggcttaagcg gctggcttcc
                                                                            1020
```

ttgtgcaaag acattggctg gcaaactaga agaggaacag atcaagaacc gagctgcatc

454

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gttacccatt gttgtctgtc atggaaaagc tgatgatgtg gtaccgttca agtttgggga
                                                                                                                                                 1140
           gaaatcttca caggctttgc tttcaaatgg gtttaagaag gtgaccttca aaccttacag
                                                                                                                                                 1200
           tgcacttggt caccacaca tcccacagga gttggatgag ttgtgcgcat ggttgacatc
           cacqctcaqc ctcqaaqqtt qatacttcct atgatqtaqc tttctqatqa aaacccttca
  5
           acticting and acting attitude attitude acting attitude acting act
           tttgtgtaag acacattcgt ctagtagcaa atcttctcct
           (2) INFORMATION FOR SEO ID NO:273:
                     (i) SEQUENCE CHARACTERISTICS:
                               (A) LENGTH: 146 amino acids
10
                               (B) TYPE: amino acid
                              (C) STRANDEDNESS:
                               (D) TOPOLOGY: linear
                   (ii) MOLECULE TYPE: peptide
                   (ix) FEATURE:
15
                               (A) NAME/KEY: peptide
                               (B) LOCATION: 1..146
                               (D) OTHER INFORMATION: / Ceres Seq. ID 1534545
                   (xi) SEQUENCE DESCRIPTION: SEQ ID NO:273:
           Lys Ile Leu Gly Leu Arg Ser Ser His Leu Lys Ile Leu Gly Ala Ser
20
                                                                                 10
           Phe Asp His Thr Leu Ser Met Ala Asp Ala Val Asn Ala Gln Thr Pro
                                                                         25
           Ser Leu Ser Glu Gln Tyr His Leu Glu Lys Glu Val Lys Gln Asp Thr
                                                                  40
25
           Ser Ala Lys Pro Val Glu Val Lys Glu Val Ala Pro Glu Val Thr Thr
                                                          5.5
           Gln Ala Glu Glu Val Lys Thr Glu Gln Ala Lys Glu Glu Ser Pro Val
                                                                                         75
                                                  70
           Glu Glu Ala Val Ser Val Val Glu Glu Lys Ser Glu Ser Ala Pro Glu
30
                                         85
                                                                                 90
           Ser Thr Glu Val Ala Ser Glu Ala Pro Ala Ala Ala Glu Asp Asn Ala
                                                                        105
                                  100
                                                                                                               110
           Glu Glu Thr Pro Ala Ala Ala Glu Glu Asn Asn Asp Glu Asn Ala Ser
                                                                120
                                                                                                     125
35
           Glu Glu Val Ala Glu Glu Thr Pro Asp Glu Ile Lys Leu Glu Thr Ala
                  130
                                                         135
                                                                                                140
           Pro Ala
           145
            (2) INFORMATION FOR SEQ ID NO:274:
40
                     (i) SEQUENCE CHARACTERISTICS:
                               (A) LENGTH: 252 amino acids
                               (B) TYPE: amino acid
                               (C) STRANDEDNESS:
                               (D) TOPOLOGY: linear
45
                   (ii) MOLECULE TYPE: peptide
                    (ix) FEATURE:
                               (A) NAME/KEY: peptide
                               (B) LOCATION: 1..252
                               (D) OTHER INFORMATION: / Ceres Seq. ID 1534546
50
                    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:274:
           Met Ser Ile Ser Gly Ala Ala Val Gly Ser Gly Arg Asn Leu Arg Arg
                                                                                10
           Ala Val Glu Phe Gly Lys Thr His Val Val Arg Pro Lys Gly Lys His
                                                                         25
                                   20
55
           Gln Ala Thr Ile Val Trp Leu His Gly Leu Gly Asp Asn Gly Ser Ser
                                                                  4.0
            Trp Ser Gln Leu Leu Glu Thr Leu Pro Leu Pro Asn Ile Lys Trp Ile
                                                          55
            Cys Pro Thr Ala Pro Ser Gln Pro Ile Ser Leu Phe Gly Gly Phe Pro
                                    70
60
                                                                                         75
            Ser Thr Ala Trp Phe Asp Val Val Asp Ile Asn Glu Asp Gly Pro Asp
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90

455 Asp Met Glu Gly Leu Asp Val Ala Ala Ala His Val Ala Asn Leu Leu 105 Ser Asn Glu Pro Ala Asp Ile Lys Leu Gly Val Gly Gly Phe Ser Met 120 125 5 Gly Ala Ala Thr Ser Leu Tyr Ser Ala Thr Cys Phe Ala Leu Gly Lys 135 140 Tyr Gly Asn Gly Asn Pro Tyr Pro Ile Asn Leu Ser Ala Ile Ile Gly 150 155 Leu Ser Gly Trp Leu Pro Cys Ala Lys Thr Leu Ala Gly Lys Leu Glu 170 175 10 165 Glu Glu Gln Ile Lys Asn Arg Ala Ala Ser Leu Pro Ile Val Val Cys 180 185 190 His Gly Lys Ala Asp Asp Val Val Pro Phe Lys Phe Gly Glu Lys Ser 195 200 205 Ser Gln Ala Leu Leu Ser Asn Gly Phe Lys Lys Val Thr Phe Lys Pro 15 210 215 220 Tyr Ser Ala Leu Gly His His Thr Ile Pro Gln Glu Leu Asp Glu Leu 225 230 235 Cys Ala Trp Leu Thr Ser Thr Leu Ser Leu Glu Gly 20 245 (2) INFORMATION FOR SEQ ID NO:275: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 155 amino acids (B) TYPE: amino acid 25 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..155 30 (D) OTHER INFORMATION: / Ceres Seq. ID 1534547 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:275: Met Glu Gly Leu Asp Val Ala Ala Ala His Val Ala Asn Leu Leu Ser 10 Asn Glu Pro Ala Asp Ile Lys Leu Gly Val Gly Gly Phe Ser Met Gly 35 25 20 Ala Ala Thr Ser Leu Tyr Ser Ala Thr Cys Phe Ala Leu Gly Lys Tyr 40 Gly Asn Gly Asn Pro Tyr Pro Ile Asn Leu Ser Ala Ile Ile Gly Leu 40 55 Ser Gly Trp Leu Pro Cys Ala Lys Thr Leu Ala Gly Lys Leu Glu Glu 70 7.5 Glu Gln Ile Lys Asn Arg Ala Ala Ser Leu Pro Ile Val Val Cys His 90 45 Gly Lys Ala Asp Asp Val Val Pro Phe Lys Phe Gly Glu Lys Ser Ser  $100 \hspace{1.5cm} 105 \hspace{1.5cm} 110$  Gln Ala Leu Leu Ser Asn Gly Phe Lys Lys Val Thr Phe Lys Pro Tyr 120 Ser Ala Leu Gly His His Thr Ile Pro Gln Glu Leu Asp Glu Leu Cys 135 50 Ala Trp Leu Thr Ser Thr Leu Ser Leu Glu Gly 150 (2) INFORMATION FOR SEQ ID NO:276: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 1592 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 60

(A) NAME/KEY: -

(B) LOCATION: 1..1592

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(D) OTHER INFORMATION: / Ceres Seq. ID 1567172
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:276:
      qtaqtctctc tttttataac cacttctcga aaactgaaac ctttgtagag agaacccata
                                                                             60
      gttcgataaa acattctttt tgcaactgag acttggcaac ttggttttac tcaaagtaag
                                                                            120
 5
      atttcttctt gttgatggtg ataagttcaa tatgatgaaa agtttggcta gtgcggttgg
                                                                            180
      agggaagacg gcgagggcat gtgatagctg cgtgaagagg cgggcacgtt ggtattgcqc
                                                                             240
      aGctgatgat gcctttcttt gccatgcttg tgacggttcg gtccactcgg caaacctct
                                                                             300
      tgctcgtaga cacgagagag ttcgcttgaa atcggctagc gccggaaaat atcgccatgc
                                                                             360
      ctcgccgcct caccaagcca cgtggcatca gggatttaca cgtaaagctc ggaccccacq
                                                                            420
10
      tqqaqqcaaq aagaqccaca cqatqqtttt tcatqatctt gtgccqgaga tqaqcacqqa
                                                                             480
      ggatcaagcg gagagttacg aggtggaaga gcagctcata tttgaggtgc cggtgatgaa
                                                                            540
      ctcgatggtt gaggagcaat gctttaacca atccctggag aaacagaatg agtttccaat
                                                                            600
      gatgccctta agtttcaaga gtagtgacga agaagatgac gacaacgctg agagttgtct
                                                                            660
      gaatggtttg ttcccaaccg acatggaact agctcagttc acagctgacg tggagactct
                                                                            720
15
      actcggtgga ggggatcgag agtttcattc catagaagaa ctagggttag gtgagatgtt
                                                                            780
      aaagatcgaa aaagaggagg tggaggaaga ggaaggagtt gtgacaagag aagtgcatga
                                                                            840
      tcaaqatgaa ggtgatgaga catccccatt tgaaataagc tttgactacg agtacacaca
                                                                            900
      caagaccaca ttcgatgaag gagaagaaga tgagaaagaa gacgtgatga agaatgtgat
                                                                            960
      qqaqatggga gtgaatgaga tgagtggtgg gattaaagaa gagaagaagg agaaggctct
                                                                           1020
20
      tatgcttaga ttggactatg aatcagtcat ttccacttgg ggaggccaag ggatcccatg
                                                                           1080
      gaccgcccgg gtgccatctg aaatagacct cgacatggtt tgtttcccaa cccataccat
                                                                           1140
      gggtgaaagt ggagcagagg ctcatcatca caaccacttc cgcggcctaG ggttacacct
                                                                           1200
      aggagatgct ggggatggag gaagagagc tagggtttca agataccgag agaaaaggag
                                                                           1260
      qacaaqqttq ttctccaaqa aqataaqqta cqaqqtacqt aaattqaatq caqataaaaq
                                                                           1320
25
      gcctcqcatq aaaqqaaqqt tcqtcaaqaq atcttcaatt gqtqttqctc actaaaqaac
                                                                           1380
      ttaattaatt atggatatta aattactttg ctctcatctt gcttttttgt tgctatagtt
                                                                           1440
      ttggtgattg ttagetttet ttttetgeat teatagagaa ttttgeacgt ttttgtgage
                                                                           1500
      tacqtatqta cataaatata tcaccaaaaa atqtqactat cttqtaaqca ctqatttata
                                                                          1560
      tagtcgatat aacgtgaatt ttgattgctg gt
30
      (2) INFORMATION FOR SEQ ID NO: 277:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 407 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
35
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..407
40
                (D) OTHER INFORMATION: / Ceres Seq. ID 1567173
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:277:
      Met Met Lys Ser Leu Ala Ser Ala Val Gly Gly Lys Thr Ala Arg Ala
                                          1.0
      Cys Asp Ser Cys Val Lys Arg Arg Ala Arg Trp Tyr Cys Ala Ala Asp
45
                                      25
      Asp Ala Phe Leu Cys His Ala Cys Asp Gly Ser Val His Ser Ala Asn
                                  40
      Pro Leu Ala Arg Arg His Glu Arg Val Arg Leu Lys Ser Ala Ser Ala
50
      Gly Lys Tyr Arg His Ala Ser Pro Pro His Gln Ala Thr Trp His Gln
                                              75
      Gly Phe Thr Arg Lys Ala Arg Thr Pro Arg Gly Gly Lys Lys Ser His
      Thr Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln
55
                                      105
      Ala Glu Ser Tyr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val
                                  120
      Met Asn Ser Met Val Glu Glu Gln Cys Phe Asn Gln Ser Leu Glu Lys
                              135
                                                 140
60
      Gln Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu
                          150
                                              155
      Glu Asp Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr
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457 170 165 Asp Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly 185 Gly Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu 5 200 Met Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Gly Val Val 215 Thr Arg Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe 230 235 10 Glu Ile Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu 250 245 Gly Glu Glu Asp Glu Lys Glu Asp Val Met Lvs Asn Val Met Glu Met 265 Gly Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys 15 280 Ala Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly 295 Gly Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu 315 320 310 20 Asp Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu 325 330 Ala His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp 345 Ala Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys 25 360 Arg Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys 370 375 Leu Asn Ala Asp Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg 390 30 Ser Ser Ile Gly Val Ala His 405 (2) INFORMATION FOR SEQ ID NO:278: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 406 amino acids 35 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 40 (A) NAME/KEY: peptide (B) LOCATION: 1..406 (D) OTHER INFORMATION: / Ceres Seq. ID 1567174 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:278: Met Lys Ser Leu Ala Ser Ala Val Gly Gly Lys Thr Ala Arg Ala Cys 45 1.0 Asp Ser Cys Val Lys Arg Arg Ala Arg Trp Tyr Cys Ala Ala Asp Asp 25 Ala Phe Leu Cys His Ala Cys Asp Gly Ser Val His Ser Ala Asn Pro 40 50 Leu Ala Arg Arg His Glu Arg Val Arg Leu Lys Ser Ala Ser Ala Gly 55 Lys Tyr Arg His Ala Ser Pro Pro His Gln Ala Thr Trp His Gln Gly 75 Phe Thr Arg Lys Ala Arg Thr Pro Arg Gly Gly Lys Lys Ser His Thr 55 85 Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln Ala 105 110 100 Glu Ser Tyr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val Met 120 125 115 60 Asn Ser Met Val Glu Glu Gln Cys Phe Asn Gln Ser Leu Glu Lys Gln 135 140 Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu Glu

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150
                                        155
     Asp Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr Asp
                   165
                       170
     Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly Gly
 5
                    185
     Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu Met
                200
     Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Gly Val Val Thr
                          215
10
     Arg Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe Glu
                       230
                                         235
     Ile Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu Gly
                   245
                                     250
     Glu Glu Asp Glu Lys Glu Asp Val Met Lys Asn Val Met Glu Met Gly
15
                                 265
     Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys Ala
                              280
     Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly Gly
                          295
20
     Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu Asp
                                        315
                      310
     Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu Ala
                                    330
                  325
     His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp Ala
25
               340 345 350
     Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys Arg
          355 360
     Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys Leu
       370 375
30
     Asn Ala Asp Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg Ser
                                        395
     Ser Ile Gly Val Ala His
                   405
     (2) INFORMATION FOR SEQ ID NO:279:
35
        (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 310 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
40
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..310
              (D) OTHER INFORMATION: / Ceres Seq. ID 1567175
45
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:279:
     Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln Ala
                                    10
     Glu Ser Tyr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val Met
                                25
                20
50
     Asn Ser Met Val Glu Glu Gln Cys Phe Asn Gln Ser Leu Glu Lys Gln
                             40
     Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu Glu
                          55
     Asp Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr Asp
55
                       70
                                         75
     Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly Gly
                                     90
     Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu Met
                                 105
     Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Glu Gly Val Val Thr
60
                           120
     Arg Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe Glu
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459 130 135 140 Ile Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu Gly 155 150 Glu Glu Asp Glu Lys Glu Asp Val Met Lys Asn Val Met Glu Met Gly 5 170 165 Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys Ala 185 Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly Gly 200 10 Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu Asp 215 Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu Ala 230 235 His His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp Ala 15 250 245 Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys Arg 260 265 Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys Leu 280 20 Asn Ala Asp Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg Ser 295 300 Ser Ile Gly Val Ala His 305 (2) INFORMATION FOR SEQ ID NO:280: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 520 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..520 (D) OTHER INFORMATION: / Ceres Seq. ID 1567555 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:280: aaaaattccg cagaattgat tatcgtcatt tcgttaagga tctctgtggg gtttgatcga 60 atttcatagc gaaggagaga gtgagagaga gattgaagca gtaatggcag gaacatctgg 120 attgctcaac gcagtgaagc caaagatcca aacgattgat attcaggccg ccgctggatg 180 gggaatcgcc gccgcagccg gtgccatctg ggtcgtccaa ccatttggtt ggataaagaa 240 40 gacattcatt gacccacccc caactgaaga gaagtgagat atcaccaaga ttcaagacta 300 agtgaagagt tttccagttt ctaagtttgt taccgcctct ttggcgtgga gctgaataaa 360 ttttggttct tttctggggc tttctggttt catttcttca atgtttgact gatttgtctt 420 ctcaaatqtt ataacaaaca agaaaagcac ttactccaag ttacttgaaa agagaaccca 480 ttttgcattt atcatgagtt attatttTcc atgtcaaagt 45 (2) INFORMATION FOR SEQ ID NO:281: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 57 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..57 55 (D) OTHER INFORMATION: / Ceres Seq. ID 1567536 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:281: Met Ala Gly Thr Ser Gly Leu Leu Asn Ala Val Lys Pro Lys Ile Gln 10 Thr Ile Asp Ile Gln Ala Ala Ala Gly Trp Gly Ile Ala Ala Ala Ala 60 25

Gly Ala Ile Trp Val Val Gln Pro Phe Gly Trp Ile Lys Lys Thr Phe 35 40 45

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Ile Asp Pro Pro Pro Thr Glu Glu Lys
          50
               55
      (2) INFORMATION FOR SEQ ID NO:282:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 765 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
10
          (ix) FEATURE:
              (A) NAME/KEY: -
                (B) LOCATION: 1..765
                (D) OTHER INFORMATION: / Ceres Seq. ID 1569689
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:282:
15
      aaccaaaaat ttcttcgtct ctgtctccaa aatcgaatca aaatctctaa agtttcaatt
     tttttgttct gttctttttt ttttttaaag aatggcttca atttctgcaa ctttgccttc
                                                                            120
      gccattgtta ctcacacaga gaaaatccaa tctcacatcg attcaaaaac tcccattttc
                                                                            180
     tctaactcga ggtacgaatg atctttctcc attatctctt actcgaaacc ctagcagcat
                                                                            240
     cagtetgatg gtgaaageta gtggagaaag etcagattea tegaetgate tegaegttgt
                                                                            300
20
     tagtacgatt cagaatgttt gggataagtc tgaagatagg ttaggtctta ttggtttggg
     ttttRgctgg tattgtagct ctttgggcat cattgaatct catcacggca attgacaaat
                                                                            420
     tgsccgttat ctcgagcgga ttcgaactag ttggtatctt gttctccacg tggttcacat
                                                                            480
      atogatatot ottgttoaaa ooggacagao aggagottto gaaaattgto aagaaatoag
                                                                            540
     tageggatat acttggeeag tgaacettgt gtgtgtgata ataeTtteat etttggaaga
                                                                          600
25
     tgatttgttt gcaagtttgt aaaattacat gacagggtgg ttgttgtttc tagtccaata
                                                                            660
     atgtcatgca tttgaaacct gtaaatactt tattgttggt ttttggttgt gagcaaaatc
                                                                           720
     aatcttttct aatttcaaag attctctttt atgattatac gtttt
      (2) INFORMATION FOR SEQ ID NO:283:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 66 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..66
                (D) OTHER INFORMATION: / Ceres Seq. ID 1569690
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:283:
40
     Asn Gln Lys Phe Leu Arg Leu Cys Leu Gln Asn Arg Ile Lys Ile Ser
                                      10
      Lys Val Ser Ile Phe Leu Phe Cys Ser Phe Phe Leu Lys Asn Gly
                                     25
      Phe Asn Phe Cys Asn Phe Ala Phe Ala Ile Val Thr His Thr Glu Lys
45
                                 40
      Ile Gln Ser His Ile Asp Ser Lys Thr Pro Ile Phe Ser Asn Ser Arg
      Tyr Glu
      65
50
      (2) INFORMATION FOR SEQ ID NO:284:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 107 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
5.5
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..107
60
                (D) OTHER INFORMATION: / Ceres Seq. ID 1569691
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:284:
      Met Ala Ser Ile Ser Ala Thr Leu Pro Ser Pro Leu Leu Leu Thr Gln
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461
                                          10
      Arg Lys Ser Asn Leu Thr Ser Ile Gln Lys Leu Pro Phe Ser Leu Thr
                                      25
      Arg Gly Thr Asn Asp Leu Ser Pro Leu Ser Leu Thr Arg Asn Pro Ser
 5
                                  40
      Ser Ile Ser Leu Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser
                             55
      Thr Asp Leu Asp Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser
                         70
                                             75
10
      Glu Asp Arg Leu Gly Leu Gly Leu Gly Phe Xaa Trp Tyr Cys Ser
                     85
                                     90
      Ser Leu Gly Ile Ile Glu Ser His His Gly Asn
                  100
      (2) INFORMATION FOR SEQ ID NO:285:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 55 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..55
                (D) OTHER INFORMATION: / Ceres Seq. ID 1569692
25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:285:
      Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser Thr Asp Leu Asp
                                          10
      Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser Glu Asp Arg Leu
                  20
                                      25
30
      Gly Leu Ile Gly Leu Gly Phe Xaa Trp Tyr Cys Ser Ser Leu Gly Ile
                              40
      Ile Glu Ser His His Gly Asn
      (2) INFORMATION FOR SEQ ID NO:286:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 782 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..782
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571042
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:286:
      atcaaccaaa acaaaacata aaaaacaagt ggaagettta aaacgagagg gagagagcaa
                                                                             60
      aaatggcgac gtcgggaacg tacgtgacgg aagttccgct aaaaggatcg gccgagaaat
                                                                            120
      actacaagag gtggaagaac gagaaccatg tetteeetga tgetategge caccacatee
                                                                            180
      aaaatgttac cgttcacgaa ggcgaacatg actctcacgg Gtctatcagg agttggaact
                                                                            240
50
      acacatggga tggaaaggag gaggtgttca aggagagaag agagatagac gatgagacca
                                                                            300
      aaacgttgac gttaagagga cttgagggtc acgtgatgga gcagctcaaa gtgtacgacg
                                                                            360
      tcgtctacca attcattccc aaatctgagg atacctgcat cggcaaaatc actttaatat
                                                                            420
      gggagaagcg caacgatgat tccccagaac caagcggcta catgaaattc gtcaagagct
                                                                            480
      tggttgctga catgggaaac cacgttagca aaacttaatc atcattccca cagtcgtcgt
                                                                            540
55
      catcatcatc atcatcatca tcatcatcat catcatcatc atcatcatca tcatcatcat
      catcatcatc atcatcacta totogattta taagttaaga tgttttcagt ataataaatg
      qqqtcttqtq qatcqttcat ttctatqtqt aaaccqtttq qttctqtatq atqcttcqat
      atattgttat gttcatgatc atatgtcggg ttcgatataa tgattcttaa gattaattta
60
     (2) INFORMATION FOR SEQ ID NO:287:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 151 amino acids
```

```
(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..151
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571043
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:287:
10
      Met Ala Thr Ser Gly Thr Tyr Val Thr Glu Val Pro Leu Lys Gly Ser
      1
                                           10
                                                               15
      Ala Glu Lys Tyr Tyr Lys Arg Trp Lys Asn Glu Asn His Val Phe Pro
                                       25
                  20
      Asp Ala Ile Gly His His Ile Gln Asn Val Thr Val His Glu Gly Glu
15
                                  40
      His Asp Ser His Gly Ser Ile Arg Ser Trp Asn Tyr Thr Trp Asp Gly
                              55
      Lys Glu Glu Val Phe Lys Glu Arg Arg Glu Ile Asp Asp Glu Thr Lys
                          70
                                              75
20
      Thr Leu Thr Leu Arg Gly Leu Glu Gly His Val Met Glu Gln Leu Lys
                                           90
      Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys Ser Glu Asp Thr Cys
                                      105
      Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg Asn Asp Asp Ser Pro
25
                                  120
              115
      Glu Pro Ser Gly Tyr Met Lys Phe Val Lys Ser Leu Val Ala Asp Met
                              135
      Gly Asn His Val Ser Lys Thr
                          150
30
      (2) INFORMATION FOR SEQ ID NO:288:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 718 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
35
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..718
40
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571079
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:288:
      attcaaagac atacaaaata attgagtttt ttttttttaa ttagaaacaa atggggttga
      gtggtgttct tcatgtggag gttgaggtta agtctccggc tgaaaagttc tgggtagccc
                                                                              120
      teggegaegg cateaatete tteeceaaag ettteeetaa egaetaeaaa aceateeaag
                                                                              180
45
      ttctagccgg cgacggcaac gctcctggct ccattcgcct cattacttat ggagaaggat
                                                                              240
      ctccactggt gaagatatcg gcggagagga tcgaagcagt ggatttggag aacaaaagca
                                                                              300
      tgtcatacag catcattggc ggcgaaatgt tggagtacta caaaacgttc aaaggaacca
                                                                              360
      tcaccgttat tcctaagaac ggtggcagcc ttctgaaatg gtctggtgag tttgagaaga
                                                                              420
      ccgcccatga gattgatgac ccacacgtca tcaaggactt tgctgtcaag aacttcaaag
                                                                              480
50
      agatagatga gtatctcctt aagcaaacta gtgcctaaca ctagaacctt taaattatat
                                                                              540
      Maagagggtt cgatcgtctc tataagattt ttctaattaa gaagttgaat aaagtggaac
                                                                              600
      ctctttatga atatccaagt ttgtgatttc ggagtttatg cagcctagta ggccataagc
                                                                              660
      tttttacaaa agccaattta gtcgaacatt ttgaaaaaat cgaacctttt ggtaaagc
      (2) INFORMATION FOR SEQ ID NO:289:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 155 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
```

(A) NAME/KEY: peptide

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(B) LOCATION: 1..155
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571080
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:289:
      Met Gly Leu Ser Gly Val Leu His Val Glu Val Glu Val Lys Ser Pro
 5
                                          10
      Ala Glu Lys Phe Trp Val Ala Leu Gly Asp Gly Ile Asn Leu Phe Pro
                                      25
      Lys Ala Phe Pro Asn Asp Tyr Lys Thr Ile Gln Val Leu Ala Gly Asp
                                  40
10
      Gly Asn Ala Pro Gly Ser Ile Arg Leu Ile Thr Tyr Gly Glu Gly Ser
                              55
      Pro Leu Val Lys Ile Ser Ala Glu Arg Ile Glu Ala Val Asp Leu Glu
                          70
                                               7.5
      Asn Lys Ser Met Ser Tyr Ser Ile Ile Gly Gly Glu Met Leu Glu Tyr
15
                                           90
      Tyr Lys Thr Phe Lys Gly Thr Ile Thr Val Ile Pro Lys Asn Gly Gly
                                      105
      Ser Leu Leu Lys Trp Ser Gly Glu Phe Glu Lys Thr Ala His Glu Ile
              115
                                  120
                                                       125
20
      Asp Asp Pro His Val Ile Lys Asp Phe Ala Val Lys Asn Phe Lys Glu
                              135
                                                   140
      Ile Asp Glu Tyr Leu Leu Lys Gln Thr Ser Ala
      145
                          150
      (2) INFORMATION FOR SEQ ID NO:290:
2.5
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 808 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..808
                (D) OTHER INFORMATION: / Ceres Seq. ID 1572097
35
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:290:
      atcaaccaaa acaaaacata aaaaacaagt ggaagcttta aaacgagagg gatagagcaa
      aaatggcgac gtcgggaacg tacgtgacgg aagttccgct aaaaggatcg gccgagaaat
                                                                             120
      actacaaqaq qtqqaaqaac qagaaccatg tcttccctga tgctatcggc caccacatcc
                                                                             180
      aaaatgttac cgttcacgaa ggcgaacatg actctcacgg gtctatcagg agttggaact
                                                                             240
40
      acacatggga tggaaaggag gaggtgttca aggagagaag agagatagac gatgagacca
                                                                             300
      aaacgttgac gttaagagga cttgagggtc acgtgatgga gcagctcaaa gtgtacgacg
                                                                             360
      togtotacca attoattocc aaatotgagg atacotgoat oggoaaaato actttaatat
                                                                             420
      gggagaagcg caacgatgat tccccagaac caagcggcta catgaaattc gtcaagagct
                                                                             480
      tggttgctga catgggaaac cacgttagCC CcAAAAAAA AAAAAaaaa cttaatcatc
                                                                             540
45
      attoccacag togtogtoat catcatcatc atcatcatca toatcatcat catcatcatc
                                                                             600
      atcatcatca tcatcatcat catcatcatc atcactatct cgatttataa gttaagatgt
                                                                             660
      tttcagtata ataaatgggg tcttgtggat cgttcatttc tatgtgtaaa ccgtttggtt
                                                                             720
      ctgtatgatg cttcgatata ttgttatgtt catgatcata tgtcgggttc gatataatga
                                                                             780
      ttcttaagat taatttacta cacatttc
50
      (2) INFORMATION FOR SEQ ID NO:291:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 195 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
55
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1...195
60
                (D) OTHER INFORMATION: / Ceres Seq. ID 1572098
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:291:
      Met Ala Thr Ser Gly Thr Tyr Val Thr Glu Val Pro Leu Lys Gly Ser
```

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464
                                     10
     Ala Glu Lys Tyr Tyr Lys Arg Trp Lys Asn Glu Asn His Val Phe Pro
               2.0
                                 25
     Asp Ala Ile Gly His His Ile Gln Asn Val Thr Val His Glu Gly Glu
5
                              40
                                                45
     His Asp Ser His Gly Ser Ile Arg Ser Trp Asn Tyr Thr Trp Asp Gly
                          55
                                           60
     Lys Glu Glu Val Phe Lys Glu Arg Arg Glu Ile Asp Asp Glu Thr Lys
                      70
                                        75
10
     Thr Leu Thr Leu Arg Gly Leu Glu Gly His Val Met Glu Gln Leu Lys
                                     90
                   85
     Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys Ser Glu Asp Thr Cys
                                 105
     Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg Asn Asp Asp Ser Pro
15
                              120
                                               125
            115
     Glu Pro Ser Gly Tyr Met Lys Phe Val Lys Ser Leu Val Ala Asp Met
                          135
     Gly Asn His Val Ser Pro Lys Lys Lys Lys Lys Leu Asn His His
                       150
                                         155
20
     165
                                     170
     180
                                  185
     Leu Asp Leu
25
            195
     (2) INFORMATION FOR SEO ID NO: 292:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 104 amino acids
              (B) TYPE: amino acid
30
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
35
              (B) LOCATION: 1..104
              (D) OTHER INFORMATION: / Ceres Seq. ID 1572099
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:292:
     Met Glu Gln Leu Lys Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys
                                     10
40
     Ser Glu Asp Thr Cys Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg
                                 25
     Asn Asp Asp Ser Pro Glu Pro Ser Gly Tyr Met Lys Phe Val Lys Ser
                             40
     Leu Val Ala Asp Met Gly Asn His Val Ser Pro Lys Lys Lys Lys
45
                         55
     Lys Leu Asn His His Ser His Ser Arg Arg His His His His His
                      70
                                        75
     85
                                     90
50
     His His His Tyr Leu Asp Leu
                100
     (2) INFORMATION FOR SEQ ID NO:293:
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 583 base pairs
55
              (B) TYPE: nucleic acid
              (C) STRANDEDNESS: single
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: DNA (genomic)
         (ix) FEATURE:
60
              (A) NAME/KEY: -
              (B) LOCATION: 1..583
              (D) OTHER INFORMATION: / Ceres Seq. ID 1572890
```

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:293:
     aaaaacaaaa ataaaaaaaa catcgcacaa gaaaataaaa gatttgtaga atcaactaag
                                                                            60
     aaaatggcta gcactatgat gactacattg cctcagttca atggtcttcg agccaccaaa
                                                                            120
     atctctgcag ctcctgtaca aggcctggca agtgttcagc ccatgagacg caagggaaat
                                                                            180
 5
     ggagetttgg gtgcaaagtg tgacttcatc ggttcatcaa caaatctgat aatggtaacg
                                                                            240
     tcgacgaccc tgattttgtt cgcggggaga ttcggacttg cgccatcagc caataggaag
                                                                            300
     gcgacagctg gacttaggtt ggaggcacgt gactcaggtc tacaaacggg tgacccggcc
                                                                            360
     gggttcacgc ttgcggacac tttggcttgt ggcaccgttg gtcatatcat cggtgtagga
                                                                            420
     gttgtccttg gccttaaaaa cattggtgct atttgaagtt cctaaagctc ttttatttgt
                                                                            480
10
     atttgtaaaa Atttgtagat ttttataaca atattctcat gcacctgaac gagatctaat
                                                                            540
     ggattttaca agtctttatg tttatcttat aatgttgtat cgc
      (2) INFORMATION FOR SEQ ID NO:294:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 151 amino acids
15
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
20
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..151
                (D) OTHER INFORMATION: / Ceres Seq. ID 1572891
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:294:
     Lys Asn Lys Asn Lys Asn Ile Ala Gln Glu Asn Lys Arg Phe Val
25
                                         10
     Glu Ser Thr Lys Lys Met Ala Ser Thr Met Met Thr Thr Leu Pro Gln
                                     25
      Phe Asn Gly Leu Arg Ala Thr Lys Ile Ser Ala Ala Pro Val Gln Gly
                                 4.0
30
      Leu Ala Ser Val Gln Pro Met Arg Arg Lys Gly Asn Gly Ala Leu Gly
                              55
     Ala Lys Cys Asp Phe Ile Gly Ser Ser Thr Asn Leu Ile Met Val Thr
     Ser Thr Thr Leu Ile Leu Phe Ala Gly Arg Phe Gly Leu Ala Pro Ser
35
                      85
                                          90
     Ala Asn Arg Lys Ala Thr Ala Gly Leu Arg Leu Glu Ala Arg Asp Ser
                                     105
                  100
      Gly Leu Gln Thr Gly Asp Pro Ala Gly Phe Thr Leu Ala Asp Thr Leu
                                 120
              115
40
      Ala Cys Gly Thr Val Gly His Ile Ile Gly Val Gly Val Leu Gly
                              135
                                                 140
      Leu Lys Asn Ile Gly Ala Ile
                          150
      (2) INFORMATION FOR SEQ ID NO:295:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 130 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..130
                (D) OTHER INFORMATION: / Ceres Seq. ID 1572892
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:295:
      Met Ala Ser Thr Met Met Thr Thr Leu Pro Gln Phe Asn Gly Leu Arg
                                          10
      Ala Thr Lys Ile Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln
                20
                                     25
60
      Pro Met Arg Arg Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe
                                 40
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Ile Gly Ser Ser Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Ile

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55
     Leu Phe Ala Gly Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala
                                           75
                        70
     Thr Ala Gly Leu Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly
 5
                                      90
                   85
     Asp Pro Ala Gly Phe Thr Leu Ala Asp Thr Leu Ala Cys Gly Thr Val
                100 105 110
     Gly His Ile Ile Gly Val Gly Val Leu Gly Leu Lys Asn Ile Gly
                               120
10
     Ala Ile
         130
     (2) INFORMATION FOR SEQ ID NO:296:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 126 amino acids
15
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
20
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..126
               (D) OTHER INFORMATION: / Ceres Seq. ID 1572893
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:296:
     Met Met Thr Thr Leu Pro Gln Phe Asn Glv Leu Arg Ala Thr Lys Ile
25
                    5
                                      10
     Ser Ala Ala Pro Val Gln Gly Leu Ala Ser Val Gln Pro Met Arg Arg
     Lys Gly Asn Gly Ala Leu Gly Ala Lys Cys Asp Phe Ile Gly Ser Ser
                               40
30
     Thr Asn Leu Ile Met Val Thr Ser Thr Thr Leu Ile Leu Phe Ala Gly
                           55
     Arg Phe Gly Leu Ala Pro Ser Ala Asn Arg Lys Ala Thr Ala Gly Leu
                        70
                                           75
     Arg Leu Glu Ala Arg Asp Ser Gly Leu Gln Thr Gly Asp Pro Ala Gly
                                      90
35
                    85
     Phe Thr Leu Ala Asp Thr Leu Ala Cys Gly Thr Val Gly His Ile Ile
                100 105
     Gly Val Gly Val Leu Gly Leu Lys Asn Ile Gly Ala Ile
             115
                               120
40
     (2) INFORMATION FOR SEQ ID NO:297:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 748 base pairs
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: single
45
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: DNA (genomic)
         (ix) FEATURE:
               (A) NAME/KEY: -
               (B) LOCATION: 1..748
50
               (D) OTHER INFORMATION: / Ceres Seq. ID 1573606
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:297:
     gaaactggtg atgttqttgg ttctttgtat cttaccggcg ataGccatgg cggcaaggag
     qagtaatatt qqaaaqaata caatggtggt tcaaggtagc acttactgtg acatttqcaa
                                                                       180
55
     atteggette gagacteetg aateeteeta etteateeee ggtgeaaegg tgaagetate
                                                                       240
     atgcaaaqac aggaaqacaa tggaaqaggt ttacacagac aaagctgtat cggacaaaqa
                                                                       300
     aggaaagtat aagttcattg tccacgacga tcacagagac cagatgtgcg atgttttgct
                                                                       360
     tgtgaaaagc tcggataaaa cctgctctaa aatctccgtt ggacgtgaga agtctcqtqt
                                                                       420
     qatcttqaac cattacagtg gcattgcctc gcagatcaga catgctaaca acatgggatt
                                                                       480
60
     cqaqaaaqaa qtqaqtqatq tqttctgctc tqctttgttt caqaaqtata tqgttgatqa
                                                                       540
     agatgaggat gatattaaaa accatctcta atctctctgt ttaatcttat gatctgctgt
                                                                       600
     tttcttcatt aatgagtttc gagttatgga agagatatat ttgtatttgt ttgattactt
                                                                       660
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atttgttgtc tttagagatg ttgactctgg tgatcggata actatctgtt tgtgtaagct 720
     tcttatatat tgatgtgtca tttccttg
     (2) INFORMATION FOR SEQ ID NO: 298:
          (i) SEQUENCE CHARACTERISTICS:
 5
               (A) LENGTH: 171 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
10
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..171
               (D) OTHER INFORMATION: / Ceres Seq. ID 1573607
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:298:
15
     Met Ala Lys Leu Val Met Leu Leu Val Leu Cys Ile Leu Pro Ala Ile
     Ala Met Ala Ala Arg Arg Ser Asn Ile Gly Lys Asn Thr Met Val Val
     Gln Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro
20
                                 40
     Glu Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys
                             55
     Asp Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp
                                             75
                         70
     Lys Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His Arg Asp Gln
25
                    85
                                        90
     Met Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys
                                    105
                                                        110
     Ile Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser
30
                                120
                                                    125
     Gly Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys
                            135 140
     Glu Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val
                        150
                                            155
35
     Asp Glu Asp Glu Asp Asp Ile Lys Asn His Leu
                    165
      (2) INFORMATION FOR SEQ ID NO:299:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 166 amino acids
40
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
45
               (A) NAME/KEY: peptide
                (B) LOCATION: 1..166
                (D) OTHER INFORMATION: / Ceres Seq. ID 1573608
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:299:
     Met Leu Val Leu Cys Ile Leu Pro Ala Ile Ala Met Ala Arq
50
                                        10
     Arg Ser Asn Ile Gly Lys Asn Thr Met Val Val Gln Gly Ser Thr Tyr
                                     25
      Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro Glu Ser Ser Tyr Phe
                                 40
55
      Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys Asp Arg Lys Thr Met
                             55
      Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp Lys Glu Gly Lys Tyr
                                             75
      Lys Phe Ile Val His Asp Asp His Arq Asp Gln Met Cys Asp Val Leu
60
                                         90
      Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys Ile Ser Val Gly Arg
                                     105
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468 Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser Gly Ile Ala Ser Gln 120 115 Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys Glu Val Ser Asp Val 135 130 140 5 Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val Asp Glu Asp Glu Asp 150 155 Asp Ile Lys Asn His Leu 165 (2) INFORMATION FOR SEQ ID NO:300: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 154 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..154 (D) OTHER INFORMATION: / Ceres Seq. ID 1573609 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:300: Met Ala Ala Arg Arg Ser Asn Ile Glv Lys Asn Thr Met Val Val Gln 10 Gly Ser Thr Tyr Cys Asp Ile Cys Lys Phe Gly Phe Glu Thr Pro Glu 20 25 25 Ser Ser Tyr Phe Ile Pro Gly Ala Thr Val Lys Leu Ser Cys Lys Asp 40 Arg Lys Thr Met Glu Glu Val Tyr Thr Asp Lys Ala Val Ser Asp Lys 55 Glu Gly Lys Tyr Lys Phe Ile Val His Asp Asp His Arg Asp Gln Met 30 Cys Asp Val Leu Leu Val Lys Ser Ser Asp Lys Thr Cys Ser Lys Ile 8.5 90 Ser Val Gly Arg Glu Lys Ser Arg Val Ile Leu Asn His Tyr Ser Gly 105 100 110 Ile Ala Ser Gln Ile Arg His Ala Asn Asn Met Gly Phe Glu Lys Glu 35 120 115 125 Val Ser Asp Val Phe Cys Ser Ala Leu Phe Gln Lys Tyr Met Val Asp 135 140 Glu Asp Glu Asp Asp Ile Lys Asn His Leu 40 150 145 (2) INFORMATION FOR SEQ ID NO:301: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 704 base pairs (B) TYPE: nucleic acid 45 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -50 (B) LOCATION: 1..704 (D) OTHER INFORMATION: / Ceres Seq. ID 1573861 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:301: aaaaaatcga atcaaaatct ctaaagtttc aatttttttg ttctgttctt ttttttttt 60 taaagaatgg cttcaatttc tgcaactttg ccttcgccat tgttactcac acagagaaaa 120 55 tocaatotoa catogattoa aaaactooca ttttototaa ctogaggtac gaatgatott 180 totocattat otottactog aaacootago agcatcagto tgatggtgaa agctagtgga 240 gaaagctcag attcatcgac tgatctcgac gttgttagta cgattcagaa tgtttgggat 300 aagtctgaaa ataggttagg tcttattggt ttgggttttg ctggtattgt agctctttgg 360 gcatcattga atctcatcac ggcaattgac aaattgcccg ttatctcgag cggattcgaa 420

ctagttggta tcttgttctc cacgtggttc acatatcgat atctcttgtt caaaccggac

agacaggagc tttcgaaaat tgtcaagaaa tcagtagcgg atatacttgg cAcagtgaac

cttqtqtqtq tqataatact tcatctttqg aaqatgattt gtttgcaagt ttqtaaaatt

480

540

600

469

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acatgacagg gtggttgttg tttctagtcc aataatgtca tgcatttgaa acctgtaaat 660
     actttattgt tggtttttgg ttgtgagcaa aatcaatctt ttct
      (2) INFORMATION FOR SEQ ID NO:302:
          (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 179 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
10
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..179
                (D) OTHER INFORMATION: / Ceres Seq. ID 1573862
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:302:
15
     Met Ala Ser Ile Ser Ala Thr Leu Pro Ser Pro Leu Leu Leu Thr Gln
                                        10
      Arg Lys Ser Asn Leu Thr Ser Ile Gln Lys Leu Pro Phe Ser Leu Thr
                                  25
              20
      Arg Gly Thr Asn Asp Leu Ser Pro Leu Ser Leu Thr Arg Asn Pro Ser
20
                                 40
      Ser Ile Ser Leu Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser
                             55
      Thr Asp Leu Asp Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser
25
      Glu Asn Arg Leu Gly Leu Ile Gly Leu Gly Phe Ala Gly Ile Val Ala
      Leu Trp Ala Ser Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val
                                    105
                                                        110
      Ile Ser Ser Gly Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe
30
                                120
      Thr Tyr Arg Tyr Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys
                                                140
                            135
      Ile Val Lys Lys Ser Val Ala Asp Ile Leu Gly Thr Val Asn Leu Val
                        150 155
      Cys Val Ile Ile Leu His Leu Trp Lys Met Ile Cys Leu Gln Val Cys
35
                                        170
                     165
      Lvs Ile Thr
      (2) INFORMATION FOR SEQ ID NO:303:
40
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 127 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..127
                (D) OTHER INFORMATION: / Ceres Seq. ID 1573863
50
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:303:
      Met Val Lys Ala Ser Gly Glu Ser Ser Asp Ser Ser Thr Asp Leu Asp
                                         10
      Val Val Ser Thr Ile Gln Asn Val Trp Asp Lys Ser Glu Asn Arg Leu
                                     25
      Gly Leu Ile Gly Leu Gly Phe Ala Gly Ile Val Ala Leu Trp Ala Ser
55
                                 40
      Leu Asn Leu Ile Thr Ala Ile Asp Lys Leu Pro Val Ile Ser Ser Gly
      Phe Glu Leu Val Gly Ile Leu Phe Ser Thr Trp Phe Thr Tyr Arg Tyr
60
                         70
                                             75
      Leu Leu Phe Lys Pro Asp Arg Gln Glu Leu Ser Lys Ile Val Lys Lys
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90

85

470 Ser Val Ala Asp Ile Leu Gly Thr Val Asn Leu Val Cys Val Ile Ile 100 105 Leu His Leu Trp Lys Met Ile Cys Leu Gln Val Cys Lys Ile Thr 115 120 125 5 (2) INFORMATION FOR SEQ ID NO:304: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 662 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..662 15 (D) OTHER INFORMATION: / Ceres Seq. ID 1574093 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:304: attttttctt tctctctc ctccctaagc aaaactaaaa caagctatgg ctggtatgct toccqqaqtt qaqtqtqcaa qqaqqcqqcq cttccacqqt qqtqctcctc cqattqaatc 120 ctcgaacaca gcttctgtgg cggctgcggc gggacacgtc tggacacggc gaccatcgtt 180 20 ctctctttac actaccaatc atgagagcca ccaggcccat gtctccttct cggagagaaq 240 tqttaqqaat aaatcttatq qaqaaqacaa cqatqaqaaa cttqacqqaq caqccaaaqa 300 ggcaaagcag aggcttaaca agcggctgag aatcccacca cgtacaagtt caggcaaaat 360 ggtaaaqaca aaqqgaataa attggagcaa ggaaagggta aacctctcgg ggacttaccq 420 accgaggtgg Gtcgggttaa agaagagccg aggaaggttg atggaatggt tcaagcggcg 480 25 agttagggaa caacaagatt gtgctatatg tctagaccgg ttcaagaagg gtgagacctt 540 gqtacaccta ccatgtqccc ataagtttca ctccatatgc ttattgcctt ggctagacac 600 taatgtttat tgcccatatt gtagaactga tatttggaat taaatgttat atttttgatg 660 (2) INFORMATION FOR SEQ ID NO:305: 30 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 198 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 35 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..198 (D) OTHER INFORMATION: / Ceres Seq. ID 1574094 40 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:305: Met Ala Gly Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Arg Phe 10 His Gly Gly Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala 25 20 45 Ala Ala Ala Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr 40 4.5 Thr Thr Asn His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arg 55 Ser Val Arg Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp 50 75 70 Gly Ala Ala Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile 90 8.5 Pro Pro Arg Thr Ser Ser Gly Lys Met Val Lys Thr Lys Gly Ile Asn 100 105 55 Trp Ser Lys Glu Arg Val Asn Leu Ser Gly Thr Tyr Arg Pro Arg Trp 120 115 125 Val Gly Leu Lys Lys Ser Arg Gly Arg Leu Met Glu Trp Phe Lys Arg 135

Arg Val Arg Glu Gln Gln Asp Cys Ala Ile Cys Leu Asp Arg Phe Lys

Lys Gly Glu Thr Leu Val His Leu Pro Cys Ala His Lys Phe His Ser

155

170

150

165

WO 00/40695 471 Ile Cys Leu Pro Trp Leu Asp Thr Asn Val Tyr Cys Pro Tyr Cys 180 185 Arg Thr Asp Ile Trp Asn 195 (2) INFORMATION FOR SEQ ID NO:306: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 195 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..195 (D) OTHER INFORMATION: / Ceres Seq. ID 1574095 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:306: Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Arg Phe His Gly Gly 10 Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala Ala Ala Ala 25 20 Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr Thr Thr Asn 4.0 His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arg Ser Val Arg 55 25 Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp Gly Ala Ala 70 Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile Pro Pro Arg 8.5 90 Thr Ser Ser Gly Lys Met Val Lys Thr Lys Gly Ile Asn Trp Ser Lys 30 105 Glu Arg Val Asn Leu Ser Gly Thr Tyr Arg Pro Arg Trp Val Gly Leu 120 Lys Lys Ser Arg Gly Arg Leu Met Glu Trp Phe Lys Arg Arg Val Arg 135 35 Glu Gln Gln Asp Cys Ala Ile Cys Leu Asp Arg Phe Lys Lys Gly Glu 150 155 Thr Leu Val His Leu Pro Cys Ala His Lys Phe His Ser Ile Cys Leu 165 170 Leu Pro Trp Leu Asp Thr Asn Val Tyr Cys Pro Tyr Cys Arg Thr Asp 40 185 Ile Trp Asn 195 (2) INFORMATION FOR SEQ ID NO:307: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 725 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 50 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..725 (D) OTHER INFORMATION: / Ceres Seq. ID 1580388 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:307: 55 ctcacacaaa aatacaacaa cttagatcag tctcaaaggg ggaaaaaaaa cttaaaagaa acattaaqaq qcaacacaaa tcacacaaaa qatcaaattq aaqcctaaqa aqaaqqcaaa

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aagtgagaag caatggctac catgttgaag gtctctcttg tattgtcatt gttaggtttt ctggtgatcg ctgtcgtgac tccatcggcg gcgaacccat tcaggaagag cgtagttctc ggagggaagt caggcgttcc aaacattcgg accaacaggg aaattcaaca acttggaagg tactgcgtgg agcaattcaa tcaacaagca cagaacgagc aaggaaacat aggatccatt gcgaaaacag acacggcaat ttcgaatcca ttgcaattta gccgagtagt gtctgctcag 420 aaacaggtcg tcgctggact caaatactat ctaaggattg aagtcactca acccaatggc 480

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tctaccagga tgtttgactc tgttgtggtt attcaaccat ggctccattc taagcagttg
                                                                             540
      ctcggtttca ctcctgttgt cagtcctgtc tactaacttt atttcttctt attcgactta
                                                                             600
      aatttccata atatgatcaa gaaaagacta aaaggtgtat gatacaaagc tattaagaat
                                                                             660
      qqqttaataq ttqqttttca tqatatqttt acqttqttca taaataaaaa caaqttqtta
                                                                             720
 5
      ttagg
      (2) INFORMATION FOR SEQ ID NO:308:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 147 amino acids
                (B) TYPE: amino acid
10
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
15
                (B) LOCATION: 1..147
                (D) OTHER INFORMATION: / Ceres Seq. ID 1580389
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:308:
      Met Ala Thr Met Leu Lys Val Ser Leu Val Leu Ser Leu Leu Gly Phe
                                         10
20
      Leu Val Ile Ala Val Val Thr Pro Ser Ala Ala Asn Pro Phe Arg Lys
                                      25
      Ser Val Val Leu Gly Gly Lys Ser Gly Val Pro Asn Ile Arg Thr Asn
                                  40
      Arg Glu Ile Gln Gln Leu Gly Arg Tyr Cys Val Glu Gln Phe Asn Gln
25
                              55
      Gln Ala Gln Asn Glu Gln Gly Asn Ile Gly Ser Ile Ala Lys Thr Asp
                          70
                                              75
      Thr Ala Ile Ser Asn Pro Leu Gln Phe Ser Arg Val Val Ser Ala Gln
30
      Lys Gln Val Val Ala Gly Leu Lys Tyr Tyr Leu Arg Ile Glu Val Thr
                                      105
      Gln Pro Asn Gly Ser Thr Arg Met Phe Asp Ser Val Val Val Ile Gln
                                 120
      Pro Trp Leu His Ser Lys Gln Leu Leu Gly Phe Thr Pro Val Val Ser
35
        130
      Pro Val Tyr
      145
      (2) INFORMATION FOR SEQ ID NO:309:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 144 amino acids
40
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
45
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..144
                (D) OTHER INFORMATION: / Ceres Seq. ID 1580390
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:309:
50
      Met Leu Lys Val Ser Leu Val Leu Ser Leu Gly Phe Leu Val Ile
                                          10
      Ala Val Val Thr Pro Ser Ala Ala Asn Pro Phe Arg Lys Ser Val Val
                  20
                                       25
      Leu Gly Gly Lys Ser Gly Val Pro Asn Ile Arg Thr Asn Arg Glu Ile
55
                                   40
      Gln Gln Leu Gly Arg Tyr Cys Val Glu Gln Phe Asn Gln Gln Ala Gln
      Asn Glu Gln Gly Asn Ile Gly Ser Ile Ala Lys Thr Asp Thr Ala Ile
                          70
60
      Ser Asn Pro Leu Gln Phe Ser Arg Val Val Ser Ala Gln Lys Gln Val
                      85
                                           90
      Val Ala Gly Leu Lys Tyr Tyr Leu Arg Ile Glu Val Thr Gln Pro Asn
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473 105 100 110 Gly Ser Thr Arg Met Phe Asp Ser Val Val Val Ile Gln Pro Trp Leu 115 120 125 His Ser Lys Gln Leu Leu Gly Phe Thr Pro Val Val Ser Pro Val Tyr 5 135 (2) INFORMATION FOR SEQ ID NO:310: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 675 base pairs (B) TYPE: nucleic acid 10 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -15 (B) LOCATION: 1..675 (D) OTHER INFORMATION: / Ceres Seq. ID 1582959 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:310: tgggcaaggc aacaacata atcaacttaa atcttatcta ctttctattt ctttttaatc aaaattaccg ttcttaacta tggcgaagtg gtttttcact atcttcttgg tttttgccct 120 20 agcctcagct ttagcttgtg gcgcaagaaa cgtcccagta ggcctctctg accaaaagaa ctacctcqqa tatqqtqqcq gatattccqq cqttqqaqac aatqqtttac cctttqqtqq 240 catcagtaga agtatateta ateceggaga taatettagat tatagagagat ttagatagatae 300 tqqtqqcqqc ttaqqcqqtq qtttqqqcqq tqqaqcaqqc aqtqqattaq qcqqtqqctt 360 aggtggtgga agtggaattg gtgccggaac cagtggagga agtaccggag gagttcattt 420 25 cccttgagtt gttactttgg tttttaaggc gtcatacggt ccttattaag ctaggtctag 480 cttaagatga tgtcataata ataatttatc atatctcttt agggttttaa actttggtat 540 tatgaattat cattagctgt ttaacgtgcg tcttaagtta ctattttaac gtatgtttga 600 atcagtctag tggcttgtcg tgtcatggct tggtccattt tcaaattcta ctttgacctt ttcgagtgtt tcacc 30 (2) INFORMATION FOR SEQ ID NO:311: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 115 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 35 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..115 40 (D) OTHER INFORMATION: / Ceres Seq. ID 1582960 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:311: Met Ala Lys Trp Phe Phe Thr Ile Phe Leu Val Phe Ala Leu Ala Ser 10 Ala Leu Ala Cys Gly Ala Arg Asn Val Pro Val Gly Leu Ser Asp Gln 45 25 Lys Asn Tyr Leu Gly Tyr Gly Gly Gly Tyr Ser Gly Val Gly Asp Asn 40 Gly Leu Pro Phe Gly Gly Val Gly Gly Val Ser Gly Pro Gly Gly 55 50 Asn Leu Gly Tyr Gly Gly Phe Gly Gly Ala Gly Gly Leu Gly Gly 70 75 Gly Leu Gly Gly Gly Ala Gly Ser Gly Leu Gly Gly Leu Gly Gly 90 Gly Ser Gly Ile Gly Ala Gly Thr Ser Gly Gly Ser Thr Gly Gly Val 55 100 His Phe Pro 115 (2) INFORMATION FOR SEQ ID NO:312: (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1076 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single

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(D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
5
                (B) LOCATION: 1..1076
                (D) OTHER INFORMATION: / Ceres Seq. ID 1663221
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:312:
     aacacgagct taccgattct cttcgtctca acatttaggg tttcagagat cggctgattt
                                                                             60
     tcacatccac aagtatggcg actcaaatca gcaagaagag aaagtttgta gcggatggtg
                                                                             120
10
     tattctacgc tgaattgaat gaggttctca caagagagct agcagaggat ggttactctq
                                                                             180
     gtgttqaggt tagggttact ccaatgagga ctgagattat catcagagct actcgtactc
                                                                             240
     aqaatqttct tqqtqaqaaq qqqaqqaqaa ttaqqqaatt qacatctctt qtccaqaaqa
                                                                             300
     gattcaagtt tccagttgac agtgttgagc tctatgctga gaaggttaac aacagaggtc
                                                                             360
      totataccat tactcagact aagtototac attacaaget totoagatat cotactatte
                                                                             420
15
     gcagggcctq ttatggtgtt ttgagatttg ttatggagag tggagctaag ggatgtgagg
                                                                             480
      tcatcqtqaq tqqaaaqctc cqtqctqcac gtqctaaqtc catqaaqttc aaqqatqqtt
                                                                             540
     acatqqtttc atctqgtcaa ccaactaagg aatacatcga tgctgcagtg agacatgttt
                                                                             600
     tgctcaqaca gggggtgttg ggaatcaagg tgaagatcat gcttgactgg gaccctacqq
                                                                             660
     gcaaatcagg accaaagaca ccattgcctg atgttgtgat cattcatgct cctaaagatg
                                                                             720
20
     atgttgtcta ctctgcacct gctcaggctg ctgctccagt tactcttgtg caaqaaqctc
                                                                             780
     cactcacaac cgtagattac cctgagatga ttcctccagt ggcctagaga agaccttttt
                                                                             840
      tactattact ccatgggatt ttgttctttt tgttataact tttctacttt tgacactctc
                                                                             900
      tagtcqtatc tcttatatcc tccaaqcaga caaqtttttt ctcatqttqt ttcttaacct
                                                                             960
      atgtcaagtt ttctggaggt tgaatgcttt tagttccatg atttttttgt tcttaagact
                                                                            1020
25
      cgtaataaga aaaagaatga cgacaacaat gcctggtatt tgatgttgtt actttq
      (2) INFORMATION FOR SEQ ID NO:313:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 250 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
35
                (B) LOCATION: 1..250
                (D) OTHER INFORMATION: / Ceres Seq. ID 1663222
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:313:
      Met Ala Thr Gln Ile Ser Lys Lys Arg Lys Phe Val Ala Asp Gly Val
                                          10
40
      Phe Tyr Ala Glu Leu Asn Glu Val Leu Thr Arg Glu Leu Ala Glu Asp
                                      25
      Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile
                                  4.0
      Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg
45
                              55
      Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys Arg Phe Lys Phe Pro
                          70
      Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val Asn Asn Arg Gly Leu
                                          90
                      8.5
50
      Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly
                  100
                                      105
      Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu
                                  120
                                                      125
      Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala
55
                              135
                                                  140
      Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Val Ser Ser
                          150
                                              155
      Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ala Ala Val Arg His Val Leu
                                          170
60
      Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Leu Asp Trp
                                      185
      Asp Pro Thr Gly Lys Ser Gly Pro Lys Thr Pro Leu Pro Asp Val Val
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475 200 205 Ile Ile His Ala Pro Lys Asp Asp Val Val Tyr Ser Ala Pro Ala Gln 220 215 Ala Ala Pro Val Thr Leu Val Gln Glu Ala Pro Leu Thr Thr Val 5 230 235 Asp Tyr Pro Glu Met Ile Pro Pro Val Ala 245 (2) INFORMATION FOR SEQ ID NO:314: (i) SEQUENCE CHARACTERISTICS: 10 (A) LENGTH: 207 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 15 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..207 (D) OTHER INFORMATION: / Ceres Seq. ID 1663223 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:314: 20 Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu 10 Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys 25 Arg Phe Lys Phe Pro Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val 25 4.0 Asn Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu 70 30 Arg Phe Val Met Glu Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser 8.5 90 Gly Lys Leu Arg Ala Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly 105 Tyr Met Val Ser Ser Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ala Ala 35 120 Val Arg His Val Leu Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys 135 Ile Met Leu Asp Trp Asp Pro Thr Gly Lys Ser Gly Pro Lys Thr Pro 150 155 40 Leu Pro Asp Val Val Ile Ile His Ala Pro Lys Asp Asp Val Val Tyr 170 165 Ser Ala Pro Ala Gln Ala Ala Pro Val Thr Leu Val Gln Glu Ala 185 180 Pro Leu Thr Thr Val Asp Tyr Pro Glu Met Ile Pro Pro Val Ala 45 195 200 (2) INFORMATION FOR SEQ ID NO:315: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 488 base pairs (B) TYPE: nucleic acid 50 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -55 (B) LOCATION: 1..488 (D) OTHER INFORMATION: / Ceres Seq. ID 1663275 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:315: gtgactaaat ttctagggtt ttgaagagtc tctctcgtcg ccgtttgtta ttcctctgtg 60 agcaaatcga cgaaatgggt cactctaatg tatggaactc tcatccgaag aagtacggtc 120 60 ctggatctcg tttatgccgt gtgtgcggga actcgcacgg tctgatccgg aagtatggtt tgaactgctg cagacagtgt ttccgtagca acgctaagga gattggattc attaagtacc

gttaatcaag caccaacttc atgattgatg cttaatgata taaacatgaa ggcgtcgatg

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ggtatqqctt ttaaqctttt qtaqtttttg aaatttttac ttttqaqaac cattqttatt
      ttgggagtta attaagttgt tgaacctcat taagcatgtc ttattttgga ttaatqatqt
                                                                             420
      tttggctatt ctcgtatttt tgttttatca gtcaaatttg actaagaaga atttgcttgt
      tattagcc
 5
      (2) INFORMATION FOR SEQ ID NO:316:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 56 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
10
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..56
15
                (D) OTHER INFORMATION: / Ceres Seq. ID 1663276
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:316:
     Met Gly His Ser Asn Val Trp Asn Ser His Pro Lys Lys Tyr Gly Pro
                                          10
      Gly Ser Arg Leu Cys Arg Val Cys Gly Asn Ser His Gly Leu Ile Arg
20
                                     25
      Lys Tyr Gly Leu Asn Cys Cys Arg Gln Cys Phe Arg Ser Asn Ala Lys
                                 40
      Glu Ile Gly Phe Ile Lys Tyr Arg
                             55
25
      (2) INFORMATION FOR SEQ ID NO:317:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 38 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..38
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1663277
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:317:
      Met Glu Leu Ser Ser Glu Glu Val Arg Ser Trp Ile Ser Phe Met Pro
                                          10
      Cys Val Arg Glu Leu Ala Arg Ser Asp Pro Glu Val Trp Phe Glu Leu
40
                  20
                                      25
      Leu Gln Thr Val Phe Pro
              35
      (2) INFORMATION FOR SEQ ID NO:318:
           (i) SEQUENCE CHARACTERISTICS:
45
                (A) LENGTH: 759 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
50
                (A) NAME/KEY: -
                (B) LOCATION: 1..759
                (D) OTHER INFORMATION: / Ceres Seq. ID 1665304
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:318:
55
      atcaaccaaa acaaaacata aaaaacaagt ggaagcttta aaacgagagg gagagagcaa
                                                                             60
      aaatgqcqac gtcgqqaacg tacgtgacgg aagttccgct aaaaggatcg qccqaqaaat
                                                                             120
      actacaagag gtggaagaac gagaaccatg tetteeetga tgetategge caccacatee
                                                                             180
      aaaatgttac cgttcacgaa ggcgaacatg actctcacgg gtctatcagg agttggaact
                                                                             240
      acacatggga tggaaaggag gaggtgttca aggagagaag agagatagac gatgagacca
                                                                             300
60
      aaacgttgac gttaagagga cttgagggtc acgtgatgga gcagctcaaa gtgtacgacg
      tegtetacea atteattece aaatetgagg atacetgeat eggeaaaate actttaatat
                                                                             420
      qggagaagcq caacqatqat tccccagaac caagcggcta catgaaattc gtcaagagct
                                                                             480
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477

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tggttgctga catgggaaac cacgttagca aaacttaatc atcattccca cagtcgtcgt
                                                                             540
      cgtcgtcatc atcatcatca tcatcatcat catcatcatc atcatcatca tcatcatcat
                                                                             600
      catcactatc tcgatttata agttaagatg ttttcagtat aataaatggg gtcttgtgga
                                                                             660
      togttoattt ctatgtgtaa accgtttggt totgtatgat gottogatat attgttatgt
                                                                             720
 5
      tcatgatcat atgtcgggtt cgatataatg attcttaag
      (2) INFORMATION FOR SEQ ID NO:319:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 151 amino acids
                (B) TYPE: amino acid
10
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
15
                (B) LOCATION: 1..151
                (D) OTHER INFORMATION: / Ceres Seq. ID 1665305
          (xi) SEQUENCE DESCRIPTION: SEO ID NO:319:
      Met Ala Thr Ser Gly Thr Tyr Val Thr Glu Val Pro Leu Lys Gly Ser
                                          10
20
      Ala Glu Lys Tyr Tyr Lys Arg Trp Lys Asn Glu Asn His Val Phe Pro
                                      25
      Asp Ala Ile Gly His His Ile Gln Asn Val Thr Val His Glu Gly Glu
                                   40
      His Asp Ser His Glv Ser Ile Arg Ser Trp Asn Tyr Thr Trp Asp Gly
25
                              55
      Lys Glu Glu Val Phe Lys Glu Arg Arg Glu Ile Asp Asp Glu Thr Lys
                          70
                                               75
      Thr Leu Thr Leu Arg Gly Leu Glu Gly His Val Met Glu Gln Leu Lys
                                           90
30
      Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys Ser Glu Asp Thr Cys
                  100
                                      105
      Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg Asn Asp Asp Ser Pro
                                   120
      Glu Pro Ser Gly Tyr Met Lys Phe Val Lys Ser Leu Val Ala Asp Met
35
                               135
                                                   140
      Gly Asn His Val Ser Lys Thr
                          150
      (2) INFORMATION FOR SEO ID NO:320:
           (i) SEQUENCE CHARACTERISTICS:
40
                 (A) LENGTH: 602 base pairs
                 (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
45
          (ix) FEATURE:
                 (A) NAME/KEY: -
                 (B) LOCATION: 1..602
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1709970
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:320:
50
      qaaqtctctt atatataaqt atcgaqaqaa ttaqqqtttt ctaqtctaqt qtttccqqqa
                                                                               60
      gcagcctgag caggaggaag aagacgaaga agaagaggag cagctactac catggttctg
                                                                              120
      qqacqqaqct ttqtcqtttc aqtqqacaqa aqatttaccc aqqaaqaqqa attaqqttta
                                                                              180
      tccgatctga ttctcaggtt ttcttgtttc ttaactcaaa atgtaagagg tacttccata
                                                                              240
      acaagttqaa gccatccaag cttgcatgga ctgccatgta cagaaagcaa cncaagaagg
                                                                              300
55
      atgcagcaca agaggctqtq aagagaagga qacqtgccac caagaagcca tactcaaggt
                                                                              360
      ccattgttgg tgctaccttg gaagtaattc agaagaagag agctgagaag cctgaagttc
                                                                              420
      gtgatgcagc cagggaagct gctctgcgtg agatcaagga aagaatcaaa aagaccaaag
                                                                              480
      atgaaaagaa ggctaagaag gtggaatttg cttctaagca acagaaggtc aaggctaatt
                                                                              540
      tececaaage tgetgetgea tecaagggte etaaggtggg aggtggtggt ggeaaaeget
                                                                              600
60
      ga
      (2) INFORMATION FOR SEQ ID NO:321:
```

(i) SEQUENCE CHARACTERISTICS:

```
478
                (A) LENGTH: 108 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..108
                (D) OTHER INFORMATION: / Ceres Seq. ID 1709971
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:321:
     Met Tyr Arg Lys Gln Xaa Lys Lys Asp Ala Ala Gln Glu Ala Val Lys
                                          10
     Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
                  20
                                      25
15
     Ala Thr Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val
                                  40
     Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
                              55
     Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Val Glu Phe Ala Ser
20
                                              75
                          70
     Lys Gln Gln Lys Val Lys Ala Asn Phe Pro Lys Ala Ala Ala Ser
                                          90
     Lys Gly Pro Lys Val Gly Gly Gly Gly Lys Arg
                  100
25
      (2) INFORMATION FOR SEQ ID NO: 322:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 59 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..59
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1709972
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:322:
      Met Gln Pro Gly Lys Leu Ceu Cys Val Arg Ser Arg Lys Glu Ser Lys
                                          10
      Arg Pro Lys Met Lys Arg Arg Leu Arg Arg Trp Asn Leu Leu Ser
40
                  20
                                      25
      Asn Arg Arg Ser Arg Leu Ile Ser Pro Lys Leu Leu His Pro Arg
                                  40
      Val Leu Arg Trp Glu Val Val Val Ala Asn Ala
                              55
45
      (2) INFORMATION FOR SEQ ID NO:323:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 726 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
50
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..726
55
                (D) OTHER INFORMATION: / Ceres Seq. ID 1711273
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:323:
      atcttcttgt ctctctcta tttcttgtta ttatccacga acgaagaaaa acctagaaaa
      cagttgaaga aagaaaatca caagagaagc catggccgga attggaccga ttactcagga
      ttgggaacca gttgtgatcc gcaagagagc tcctaacgct gcagctaagc gcgacgagaa
60
      gactgtcaac gccgctcgtc gaagcggcgc cgatattgag accgttcgaa aattcaatgc
```

tggatcgaac aaggctgcat caagcggcac ctccttgaac acaaagaagc tagatgatga

tactgagaac ttatctcatg atcgtgtgcc cactgaattg aagaaagcca tcatgcaagc

60

120

180

240

300

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tagaggggag aasaagctga ctcagtccca acttgcccat ctgatcaatg agaagccaca
                                                                           420
     agtgatccaa gaatacgagt cggggaaagc aattccgaat caacagatcc tttcaaaqct
                                                                           480
      ggagagggca cttggtgcta aactccgtgg aaagaagtag aagtttagaa caaagctctt
                                                                           540
      aaaggtaaca aaaagctgat cgcagtttct ctccagtcca catgctttac catatcctaa
                                                                           600
 5
     aaactatatc tatgtatggt ttggtttaat ggcgtagtag tttgttgcga ggaatctttc
                                                                           660
      720
      ttcttt
      (2) INFORMATION FOR SEQ ID NO:324:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 142 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
15
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..142
                (D) OTHER INFORMATION: / Ceres Seq. ID 1711274
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:324:
20
     Met Ala Gly Ile Gly Pro Ile Thr Gln Asp Trp Glu Pro Val Val Ile
                                         1.0
      Arg Lys Arg Ala Pro Asn Ala Ala Lys Arg Asp Glu Lys Thr Val
                                     25
      Asn Ala Ala Arg Arg Ser Gly Ala Asp Ile Glu Thr Val Arg Lys Phe
25
                                 40
      Asn Ala Gly Ser Asn Lys Ala Ala Ser Ser Gly Thr Ser Leu Asn Thr
                             55
      Lys Lys Leu Asp Asp Asp Thr Glu Asn Leu Ser His Asp Arg Val Pro
                         70
                                             75
30
      Thr Glu Leu Lys Lys Ala Ile Met Gln Ala Arg Gly Glu Xaa Lys Leu
                                         90
                     8.5
      Thr Gln Ser Gln Leu Ala His Leu Ile Asn Glu Lys Pro Gln Val Ile
                                     105
                  100
                                                         110
      Gln Glu Tyr Glu Ser Gly Lys Ala Ile Pro Asn Gln Gln Ile Leu Ser
                                 120
35
                                                     125
             115
      Lys Leu Glu Arg Ala Leu Gly Ala Lys Leu Arg Gly Lys Lys
                             135
      (2) INFORMATION FOR SEQ ID NO:325:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 686 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
45
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..686
                (D) OTHER INFORMATION: / Ceres Seq. ID 1715423
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:325:
50
      aacaataaaa caaagtgatt tattatggcg ttaagagaga tgataataga tgagagtgat
                                                                            60
      qacataqaqa qctacaqtqa tcaatctctt tqtcttqata aaqccaaqqa qcttcttqct
                                                                           120
      ctcatcaaac tacccacqqq attqttqcca ttqaaqqata tqacqqaaqt tqqqtacaac
      aaaacaaaaq ggtttqtqtg gatqaggctg agaaqcaaga ttgagcatac attccqtqaq
                                                                           240
      ataqqtcqca qaqttttata tqacacqqYa qataactqcg tttqttqaqq accqccqtat
                                                                           300
55
      qaqqaqactt accqqaqtca aaaqcaaaga qctcatqatc tggqttcctg tqaatqatat
                                                                           360
      cttcatcaaa qaqaaaqatc ctqaqaaqat cacctttqct aataccaccq qcctqtcacq
                                                                           420
      aacatttaaa gtttcaqcat ttcaatgtga aggttgataa ataaaaagaa aaqaqqaqta
                                                                           480
      accaqacttc cttgatqaaa gcaaccaata aagaacagaa gttgttccat actcaataaq
                                                                           540
      agatagttta attaaatatc aaggacacgc ctttcttaca atgaattgct tatggatcat
                                                                           600
60
      cacaqtaaat aaacaqqttc tqcaattqat qccttqatqt ataacagtga gtttccaatc
                                                                           660
      aacatattag aatcgattca ttggtt
      (2) INFORMATION FOR SEQ ID NO: 326:
```

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(i) SEOUENCE CHARACTERISTICS:
               (A) LENGTH: 87 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
 5
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..87
10
               (D) OTHER INFORMATION: / Ceres Seq. ID 1715424
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:326:
     Met Ala Leu Arg Glu Met Ile Ile Asp Glu Ser Asp Asp Ile Glu Ser
                          10
     Tyr Ser Asp Gln Ser Leu Cys Leu Asp Lys Ala Lys Glu Leu Leu Ala
15
                                25
     Leu Ile Lys Leu Pro Thr Gly Leu Leu Pro Leu Lys Asp Met Thr Glu
                                40
     Val Gly Tyr Asn Lys Thr Lys Gly Phe Val Trp Met Arg Leu Arg Ser
                         55
20
     Lys Ile Glu His Thr Phe Arg Glu Ile Gly Arg Arg Val Leu Tyr Asp
                        70
     Thr Xaa Asp Asn Cys Val Cys
                 85
      (2) INFORMATION FOR SEQ ID NO:327:
25
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 82 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
30
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..82
               (D) OTHER INFORMATION: / Ceres Seq. ID 1715425
35
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:327:
     Met Ile Ile Asp Glu Ser Asp Asp Ile Glu Ser Tyr Ser Asp Gln Ser
                                       10
     Leu Cys Leu Asp Lys Ala Lys Glu Leu Leu Ala Leu Ile Lys Leu Pro
                      25
40
      Thr Gly Leu Pro Leu Lys Asp Met Thr Glu Val Gly Tyr Asn Lys
                                40
      Thr Lys Gly Phe Val Trp Met Arg Leu Arg Ser Lys Ile Glu His Thr
                         55
                                  60
      Phe Arg Glu Ile Gly Arg Arg Val Leu Tyr Asp Thr Xaa Asp Asn Cys
45
                                            75
      Val Cys
      (2) INFORMATION FOR SEQ ID NO:328:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 65 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 1715426
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:328:
     Met Thr Arg Xaa Ile Thr Ala Phe Val Glu Asp Arg Arg Met Arg Arg
60
                                        10
      Leu Thr Gly Val Lys Ser Lys Glu Leu Met Ile Trp Val Pro Val Asn
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481
                                      25
     Asp Ile Phe Ile Lys Glu Lys Asp Pro Glu Lys Ile Thr Phe Ala Asn
                                  40
                                                      45
     Thr Thr Gly Leu Ser Arg Thr Phe Lys Val Ser Ala Phe Gln Cys Glu
5
                              55
     Gly
      65
      (2) INFORMATION FOR SEQ ID NO:329:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 829 base pairs
               (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
15
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..829
                (D) OTHER INFORMATION: / Ceres Seq. ID 1715962
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:329:
20
     aaaaacgccg tcgagaaatc tccctctagg gtttctttga tcaaaccaga gagccgtcag
                                                                             60
     ggagagcaaa acaaagatcg gagaagatga agacgattct ttcttccgaa acgatggaca
                                                                            120
      tccccgacag tgttaccatc aaggttcacg ctaaagtgat cgaagtcgaa ggacctcgcg
                                                                            180
      ggaagettgt tegegattte aageatetea acetegattt ceagetgate aaggateeag
                                                                            240
     agactggaaa gaaqaagctt aagatcgatt cgtggtttgg aacacgcaaa accagcqcct
                                                                            300
25
     ccatcagaac cqctcttaqc cacqtcqata acttgatctc cqqtqttacc aqaqqtttcc
                                                                            360
      gttacaagat gaggttegtg tacgeceatt tteceateaa egeeteeate ggeggtgaeg
                                                                            420
      gaaagtctat cgagatccgt aacttccttg gcgagaagaa ggtgaggaag gtagagatgt
                                                                            480
      tggatggtqt aacCattgtt cgaYctgaga aggtgaagga tgagattgtt cttgacggta
                                                                            540
      acgacatcqa qcttqtttca aqqtcatqcq ctttqatcaa ccaqaaatqt cacqtqaaqa
                                                                            600
30
      agaaggatat caggaagttt cttgatggta tctatgttag cgagaaaagc aagatcgtag
                                                                            660
      aggaagaatg aatgcccata tcatcgtctt agctcctatt tcttgtcttt tgaatgttag
                                                                            720
      tttcgtttta tggaccaaat ccaccggtgt tgcaaaactc tggctatccc ttgttccctt
                                                                            780
      tttgtgttcg agattttatt tacttatgca caagttttgg agacgagct
      (2) INFORMATION FOR SEQ ID NO:330:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 194 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..194
                (D) OTHER INFORMATION: / Ceres Seq. ID 1715963
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:330:
      Met Lys Thr Ile Leu Ser Ser Glu Thr Met Asp Ile Pro Asp Ser Val
                                          10
      Thr Ile Lys Val His Ala Lys Val Ile Glu Val Glu Gly Pro Arg Gly
                  20
                                      25
50
      Lys Leu Val Arg Asp Phe Lys His Leu Asn Leu Asp Phe Gln Leu Ile
                                  40
      Lys Asp Pro Glu Thr Gly Lys Lys Leu Lys Ile Asp Ser Trp Phe
      Gly Thr Arg Lys Thr Ser Ala Ser Ile Arg Thr Ala Leu Ser His Val
55
      Asp Asn Leu Ile Ser Gly Val Thr Arg Gly Phe Arg Tyr Lys Met Arg
      Phe Val Tyr Ala His Phe Pro Ile Asn Ala Ser Ile Gly Gly Asp Gly
                                     105
                  100
60
      Lys Ser Ile Glu Ile Arg Asn Phe Leu Gly Glu Lys Lys Val Arg Lys
              115
                                  120
                                                      125
      Val Glu Met Leu Asp Gly Val Thr Ile Val Arg Xaa Glu Lys Val Lys
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135
                                           140
     Asp Glu Ile Val Leu Asp Gly Asn Asp Ile Glu Leu Val Ser Arg Ser
            150 155 160
     Cys Ala Leu Ile Asn Gln Lys Cys His Val Lys Lys Asp Ile Arg
5
                          170 175
     Lys Phe Leu Asp Gly Ile Tyr Val Ser Glu Lys Ser Lys Ile Val Glu
                                185
     Glu Glu
10
     (2) INFORMATION FOR SEQ ID NO:331:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 185 amino acids
              (B) TYPE: amino acid
             (C) STRANDEDNESS:
15
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..185
20
             (D) OTHER INFORMATION: / Ceres Seq. ID 1715964
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:331:
     Met Asp Ile Pro Asp Ser Val Thr Ile Lys Val His Ala Lys Val Ile
               5 10
     Glu Val Glu Gly Pro Arg Gly Lys Leu Val Arg Asp Phe Lys His Leu
25
                                2.5
     Asn Leu Asp Phe Gln Leu Ile Lys Asp Pro Glu Thr Gly Lys Lys
                             40
     Leu Lys Ile Asp Ser Trp Phe Gly Thr Arg Lys Thr Ser Ala Ser Ile
                         5.5
30
     Arg Thr Ala Leu Ser His Val Asp Asn Leu Ile Ser Gly Val Thr Arg
                                       75
                      70
     Gly Phe Arg Tyr Lys Met Arg Phe Val Tyr Ala His Phe Pro Ile Asn
                                 90 95
                  85
     Ala Ser Ile Gly Gly Asp Gly Lys Ser Ile Glu Ile Arg Asn Phe Leu
35
                                105
     Gly Glu Lys Lys Val Arg Lys Val Glu Met Leu Asp Gly Val Thr Ile
                            120 125
     Val Arg Xaa Glu Lys Val Lys Asp Glu Ile Val Leu Asp Gly Asn Asp
                         135 140
40
     Ile Glu Leu Val Ser Arg Ser Cys Ala Leu Ile Asn Gln Lys Cys His
                    150 155 160
     Val Lys Lys Lys Asp Ile Arg Lys Phe Leu Asp Gly Ile Tyr Val Ser
                165 170
     Glu Lys Ser Lys Ile Val Glu Glu Glu
45
            180 185
     (2) INFORMATION FOR SEQ ID NO:332:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 100 amino acids
              (B) TYPE: amino acid
50
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
             (A) NAME/KEY: peptide
55
              (B) LOCATION: 1..100
              (D) OTHER INFORMATION: / Ceres Seq. ID 1715965
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:332:
     Met Arg Phe Val Tyr Ala His Phe Pro Ile Asn Ala Ser Ile Gly Gly
                             10
     Asp Gly Lys Ser Ile Glu Ile Arg Asn Phe Leu Gly Glu Lys Lys Val
60
                                25
     Arg Lys Val Glu Met Leu Asp Gly Val Thr Ile Val Arg Xaa Glu Lys
```

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483
                                 40
                                                      45
     Val Lys Asp Glu Ile Val Leu Asp Gly Asn Asp Ile Glu Leu Val Ser
                             5.5
     Arg Ser Cys Ala Leu Ile Asn Gln Lys Cys His Val Lys Lys Lys Asp
 5
                         70
                                             75
     Ile Arg Lys Phe Leu Asp Gly Ile Tyr Val Ser Glu Lys Ser Lys Ile
     Val Glu Glu Glu
                  100
10
     (2) INFORMATION FOR SEQ ID NO:333:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 675 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
15
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..675
20
                (D) OTHER INFORMATION: / Ceres Seq. ID 1808584
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:333:
     aacttgtcac ctgcagagaa agaaagaagc cctagatttt gtcaaaaggc ggttgcagaa
     caaaaaacca tggccggaat tggaccgata actcaggatt gggagccggt ggtgatccgt
                                                                            120
     aagaaacccg ctaacgccgc tgccaagcgc gacgagaaaa ctgtcaacgc cgctcgtcga
                                                                            180
25
     teeggegeeg atategagae egteagaaaa tteaatgetg gaaccaacaa ggeggeatea
                                                                            240
     agcggcacat ctctgaacac aaaaatgctt gatgatgaca ctgagaacct tactcatgaa
                                                                            300
     cgtgtgccta ctgagctaaa gaaagccatt atgcaagcca ggacagacaa gaagctaacc
                                                                            360
     cagtcccaac ttgctcaaat catcaatgag aagccacaag tgattcaaga gtatgagtct
                                                                           420
     ggcaaagcta tacccaacca gcaaatcctt tctaagctgg agagagcgct tggagctaag
                                                                           480
30
     cttcgtggaa agaagtgagc caagttctac tgatgtagca agtaacaaga atcaatgctt
                                                                           540
     togtotaatq cogtaacttt gocaagaaga atattttotg attgtaagaa agcaaaaccq
                                                                          600
     tttgaatgtt tgtttcgttg atggaatctc tatctcataa actcatatca atataataac
                                                                           660
     ttgggtcttt tcatc
      (2) INFORMATION FOR SEQ ID NO:334:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 165 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
40
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..165
                (D) OTHER INFORMATION: / Ceres Seq. ID 1808585
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:334:
     Asn Leu Ser Pro Ala Glu Lys Glu Arg Ser Pro Arg Phe Cys Gln Lys
                                         10
     Ala Val Ala Glu Gln Lys Thr Met Ala Gly Ile Gly Pro Ile Thr Gln
50
      Asp Trp Glu Pro Val Val Ile Arg Lys Lys Pro Ala Asn Ala Ala
                                  4.0
      Lys Arg Asp Glu Lys Thr Val Asn Ala Arg Arg Ser Gly Ala Asp
                              55
      Ile Glu Thr Val Arg Lys Phe Asn Ala Gly Thr Asn Lys Ala Ala Ser
5.5
      Ser Gly Thr Ser Leu Asn Thr Lys Met Leu Asp Asp Asp Thr Glu Asn
      Leu Thr His Glu Arg Val Pro Thr Glu Leu Lys Lys Ala Ile Met Gln
                  100
                                     105
                                                         110
60
      Ala Arg Thr Asp Lys Lys Leu Thr Gln Ser Gln Leu Ala Gln Ile Ile
                                 120
```

Asn Glu Lys Pro Gln Val Ile Gln Glu Tyr Glu Ser Gly Lys Ala Ile

```
135
                                                140
      Pro Asn Gln Gln Ile Leu Ser Lys Leu Glu Arg Ala Leu Gly Ala Lys
                        150
                             155
     Leu Arg Gly Lys Lys
 5
                     165
      (2) INFORMATION FOR SEQ ID NO:335:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 142 amino acids
               (B) TYPE: amino acid
10
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
15
               (B) LOCATION: 1..142
               (D) OTHER INFORMATION: / Ceres Seg. ID 1808586
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:335:
     Met Ala Gly Ile Gly Pro Ile Thr Gln Asp Trp Glu Pro Val Val Ile
                                       10
     Arg Lys Lys Pro Ala Asn Ala Ala Ala Lys Arg Asp Glu Lys Thr Val
20
                 20
                                    25
     Asn Ala Ala Arg Arg Ser Gly Ala Asp Ile Glu Thr Val Arg Lys Phe
                                40
     Asn Ala Gly Thr Asn Lys Ala Ala Ser Ser Gly Thr Ser Leu Asn Thr
25
                            55
     Lys Met Leu Asp Asp Asp Thr Glu Asn Leu Thr His Glu Arg Val Pro
                       70
     Thr Glu Leu Lys Lys Ala Ile Met Gln Ala Arg Thr Asp Lys Lys Leu
                                     90
                     85
30
     Thr Gln Ser Gln Leu Ala Gln Ile Ile Asn Glu Lys Pro Gln Val Ile
                                    105
     Gln Glu Tyr Glu Ser Gly Lys Ala Ile Pro Asn Gln Gln Ile Leu Ser
                         120
     Lys Leu Glu Arg Ala Leu Gly Ala Lys Leu Arg Gly Lys Lys
35
                             135
      (2) INFORMATION FOR SEQ ID NO:336:
          (i) SEOUENCE CHARACTERISTICS:
               (A) LENGTH: 630 base pairs
               (B) TYPE: nucleic acid
40
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: DNA (genomic)
         (ix) FEATURE:
               (A) NAME/KEY: -
45
               (B) LOCATION: 1..630
               (D) OTHER INFORMATION: / Ceres Seq. ID 1808591
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:336:
     aacactacaa caattttcat acctcatatt ctcttccatg gcaaccacca agcacctcct
                                                                          60
     cctcttcctc ctcatcatcg ctatcatcac cattacatct tccacttcac ttccatttct
                                                                          120
50
     cacgacggag caaaaccaaa tcgcaaccaa aatcatagac gcaatggtct caagtggctc
                                                                         180
     tttcgaagat tggagcggag cgttcctcaa caacaacgac gaattaaacg gtccagtcct
                                                                          240
     aacttcaact ctcttcctcc ccaaaacatc cgtggaagga atcaacgcca cgtcaccact
                                                                          300
     agttgcttct taccatattg ttccacaatg gcttgacttc tccgtcataa gtctcatgat
                                                                          360
     gcctttctct cgcatcccta cactcctctc tggacactct atagtcgtta ccaacaattc
55
     agettetggt titactettg atggtgttet catetetgag ceagatttat tegtetetee
     tactatagtt atccatcgaa tggcttttcc atttaacttc tctcgttacg gtggtggcga
                                                                         540
     tatatagtta tttacttttg attcgcttct ccactttgtt tctttctact tttactaata
                                                                          600
     cactttcaaa taattaacaa cgaacttgac
     (2) INFORMATION FOR SEQ ID NO:337:
60
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 181 amino acids
               (B) TYPE: amino acid
```

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485
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
5
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..181
             (D) OTHER INFORMATION: / Ceres Seq. ID 1808592
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:337:
     Thr Leu Gln Gln Phe Ser Tyr Leu Ile Phe Ser Ser Met Ala Thr Thr
                                   10
10
     Lys His Leu Leu Phe Leu Leu Ile Ile Ala Ile Ile Thr Ile Thr
          20
                   25
     Ser Ser Thr Ser Leu Pro Phe Leu Thr Thr Glu Gln Asn Gln Ile Ala
                40
15
     Thr Lys Ile Ile Asp Ala Met Val Ser Ser Gly Ser Phe Glu Asp Trp
      50 55
                                60
     Ser Gly Ala Phe Leu Asn Asn Asn Glu Leu Asn Gly Pro Val Leu
            70 ·
     Thr Ser Thr Leu Phe Leu Pro Lys Thr Ser Val Glu Gly Ile Asn Ala
20
                                   90
     Thr Ser Pro Leu Val Ala Ser Tyr His Ile Val Pro Gln Trp Leu Asp
                               105
     Phe Ser Val Ile Ser Leu Met Met Pro Phe Ser Arg Ile Pro Thr Leu
           115 120
25
     Leu Ser Gly His Ser Ile Val Val Thr Asn Asn Ser Ala Ser Gly Phe
       130 135
                                 140
     Thr Leu Asp Gly Val Leu Ile Ser Glu Pro Asp Leu Phe Val Ser Pro
        150 155 160
     Thr Ile Val Ile His Arg Met Ala Phe Pro Phe Asn Phe Ser Arg Tyr
30
               165 170
     Gly Gly Gly Asp Ile
               180
     (2) INFORMATION FOR SEQ ID NO:338:
         (i) SEQUENCE CHARACTERISTICS:
35
             (A) LENGTH: 169 amino acids
             (B) TYPE: amino acid
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
40
        (ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..169
             (D) OTHER INFORMATION: / Ceres Seq. ID 1808593
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:338:
45
     Met Ala Thr Thr Lys His Leu Leu Phe Leu Leu Ile Ile Ala Ile
            5
                      10
     Ile Thr Ile Thr Ser Ser Thr Ser Leu Pro Phe Leu Thr Thr Glu Gln
                                25
     Asn Gln Ile Ala Thr Lys Ile Ile Asp Ala Met Val Ser Ser Gly Ser
50
                            40
     Phe Glu Asp Trp Ser Gly Ala Phe Leu Asn Asn Asp Glu Leu Asn
                         5.5
     Gly Pro Val Leu Thr Ser Thr Leu Phe Leu Pro Lys Thr Ser Val Glu
         70
                                      75
55
     Gly Ile Asn Ala Thr Ser Pro Leu Val Ala Ser Tyr His Ile Val Pro
                                  90
                  85
     Gln Trp Leu Asp Phe Ser Val Ile Ser Leu Met Met Pro Phe Ser Arg
              100 105
     Ile Pro Thr Leu Leu Ser Gly His Ser Ile Val Val Thr Asn Asn Ser
60
      115 120 125
     Ala Ser Gly Phe Thr Leu Asp Gly Val Leu Ile Ser Glu Pro Asp Leu
```

135

140

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Phe Val Ser Pro Thr Ile Val Ile His Arg Met Ala Phe Pro Phe Asn
                          150
                                               155
      Phe Ser Arg Tyr Gly Gly Gly Asp Ile
                      165
 5
      (2) INFORMATION FOR SEQ ID NO:339:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 127 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
10
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..127
15
                (D) OTHER INFORMATION: / Ceres Seq. ID 1808594
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:339:
      Met Val Ser Ser Gly Ser Phe Glu Asp Trp Ser Gly Ala Phe Leu Asn
                                          10
      Asn Asn Asp Glu Leu Asn Gly Pro Val Leu Thr Ser Thr Leu Phe Leu
20
                  20
                                      25
      Pro Lys Thr Ser Val Glu Gly Ile Asn Ala Thr Ser Pro Leu Val Ala
                                  40
      Ser Tyr His Ile Val Pro Gln Trp Leu Asp Phe Ser Val Ile Ser Leu
                              55
25
      Met Met Pro Phe Ser Arg Ile Pro Thr Leu Leu Ser Gly His Ser Ile
                          70
                                               75
      Val Val Thr Asn Asn Ser Ala Ser Gly Phe Thr Leu Asp Gly Val Leu
                                          90
                      8.5
      Ile Ser Glu Pro Asp Leu Phe Val Ser Pro Thr Ile Val Ile His Arg
30
                  100
                                      105
      Met Ala Phe Pro Phe Asn Phe Ser Arg Tyr Gly Gly Gly Asp Ile
              115
                                  120
      (2) INFORMATION FOR SEQ ID NO:340:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 717 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
40
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..717
                (D) OTHER INFORMATION: / Ceres Seq. ID 1920563
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:340:
45
      cttgtttata ctaatgacca aggagagact gttaaaggag tttgctcaaa tttcttgtgt
                                                                              60
      gatttggcac ccggaagtga tgttaagctg actggtcctg taggcaaaga aatgcttatg
                                                                             120
      cctaaggatc caaacgccac cgtaattatg cttgccacag ggacaggaat tgctcctttc
                                                                             180
      aggtctttct tatggaagat gttctttgag aaacatgatg actacaagtt caatggctta
                                                                             240
      gcttggttgt tcttgggtgt accaaccact agctcattgc tctaccaaga ggagtttgat
                                                                             300
      aagatgaaag caaaggcccc cgagaacttc agggtggatt acgcgataag cagagaacaa
50
                                                                             360
      gcgaacgata aaggagagaa aatgtatatc cagactcgga tggcacagta cgcagctgaa
                                                                             420
      ttatgggagt tgttgaagaa agacaacact tttgtttaca tgtgtggact caagggaatg
                                                                             480
      gagaaaggaa ttgatgacat tatggtctca ttggctgcaa atgacggtat tgactggttt
                                                                             540
      gattacaaga agcagttgaa gaaggcagag caatggaacg ttgaagtcta ctgatcaaaa
                                                                             600
55
      agcetttgae atttetgtag caaagtatag etgaacaaaa etgtaatttt egetteegaa
                                                                             660
      tttctgtatt ttgaagataa gttttttaga tatgttatac taaaaaaaga gttcttt
      (2) INFORMATION FOR SEQ ID NO:341:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 197 amino acids
60
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
```

487

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(ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..197
 5
              (D) OTHER INFORMATION: / Ceres Seq. ID 1920564
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:341:
     Leu Val Tyr Thr Asn Asp Gln Gly Glu Thr Val Lys Gly Val Cys Ser
                                      10
     Asn Phe Leu Cys Asp Leu Ala Pro Gly Ser Asp Val Lys Leu Thr Gly
10
                                 25
     Pro Val Gly Lys Glu Met Leu Met Pro Lys Asp Pro Asn Ala Thr Val
                              40
     Ile Met Leu Ala Thr Gly Thr Gly Ile Ala Pro Phe Arg Ser Phe Leu
                           55
15
     Trp Lys Met Phe Phe Glu Lys His Asp Asp Tyr Lys Phe Asn Gly Leu
                             75
     Ala Trp Leu Phe Leu Gly Val Pro Thr Thr Ser Ser Leu Leu Tyr Gln
                                      90
     Glu Glu Phe Asp Lys Met Lys Ala Lys Ala Pro Glu Asn Phe Arg Val
20
                100
                                  105
     Asp Tyr Ala Ile Ser Arg Glu Gln Ala Asn Asp Lys Gly Glu Lys Met
                              120
     Tyr Ile Gln Thr Arg Met Ala Gln Tyr Ala Ala Glu Leu Trp Glu Leu
                          135 140
25
     Leu Lys Lys Asp Asn Thr Phe Val Tyr Met Cys Gly Leu Lys Gly Met
     145 150 155
     Glu Lys Gly Ile Asp Asp Ile Met Val Ser Leu Ala Ala Asn Asp Gly
                  165 170 175
     Ile Asp Trp Phe Asp Tyr Lys Lys Gln Leu Lys Lys Ala Glu Gln Trp
30
     180 185
Asn Val Glu Val Tyr
            195
     (2) INFORMATION FOR SEQ ID NO:342:
          (i) SEQJENCE CHARACTERISTICS:
35
              (A) LENGTH: 160 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
40
        (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..160
              (D) OTHER INFORMATION: / Ceres Seq. ID 1920565
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:342:
45
     Met Leu Met Pro Lys Asp Pro Asn Ala Thr Val Ile Met Leu Ala Thr
                                     10
     Gly Thr Gly Ile Ala Pro Phe Arg Ser Phe Leu Trp Lys Met Phe Phe
                                  25
     Glu Lys His Asp Asp Tyr Lys Phe Asn Gly Leu Ala Trp Leu Phe Leu
50
                              40
     Gly Val Pro Thr Thr Ser Ser Leu Leu Tyr Gln Glu Glu Phe Asp Lys
                           55
     Met Lys Ala Lys Ala Pro Glu Asn Phe Arg Val Asp Tyr Ala Ile Ser
                       70
     Arg Glu Gln Ala Asn Asp Lys Gly Glu Lys Met Tyr Ile Gln Thr Arg
55
                   85
     Met Ala Gln Tyr Ala Ala Glu Leu Trp Glu Leu Leu Lys Lys Asp Asn
                                 105
               100
     Thr Phe Val Tyr Met Cys Gly Leu Lys Gly Met Glu Lys Gly Ile Asp
60
       115 120
     Asp Ile Met Val Ser Leu Ala Ala Asn Asp Gly Ile Asp Trp Phe Asp
```

135

130

```
Tyr Lys Lys Gln Leu Lys Lys Ala Glu Gln Trp Asn Val Glu Val Tyr
                          150
                                               155
      (2) INFORMATION FOR SEQ ID NO:343:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 158 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
10
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..158
                (D) OTHER INFORMATION: / Ceres Seq. ID 1920566
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:343:
15
      Met Pro Lys Asp Pro Asn Ala Thr Val Ile Met Leu Ala Thr Gly Thr
                                          10
      Gly Ile Ala Pro Phe Arg Ser Phe Leu Trp Lys Met Phe Phe Glu Lys
                                      25
      His Asp Asp Tyr Lys Phe Asn Gly Leu Ala Trp Leu Phe Leu Gly Val
20
                                  40
      Pro Thr Thr Ser Ser Leu Leu Tyr Gln Glu Glu Phe Asp Lys Met Lys
      Ala Lys Ala Pro Glu Asn Phe Arg Val Asp Tyr Ala Ile Ser Arg Glu
                          70
25
      Gln Ala Asn Asp Lys Gly Glu Lys Met Tyr Ile Gln Thr Arg Met Ala
                                          90
      Gln Tyr Ala Ala Glu Leu Trp Glu Leu Lys Lys Asp Asn Thr Phe
                                      105
      Val Tyr Met Cys Gly Leu Lys Gly Met Glu Lys Gly Ile Asp Asp Ile
30
              115
                                  120
                                                      125
      Met Val Ser Leu Ala Ala Asn Asp Gly Ile Asp Trp Phe Asp Tyr Lys
                              135
                                                  1/0
      Lys Gln Leu Lys Lys Ala Glu Gln Trp Asn Val Gl. Val Tyr
                          150
                                              155
35
      (2) INFORMATION FOR SEQ ID NO:344:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 2192 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
40
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..2192
45
                (D) OTHER INFORMATION: / Ceres Seq. ID 1974419
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:344:
      aattcaagtc atctctatcc gtcaacaata caaaccaacc ttctcaattc ctctcttttc
                                                                             60
      atccacattt tcactagcta atggcttcta gttcaatgag cacctcatca tggacagcta
                                                                             120
      gagaggacaa gcaattcgaa atggcgttgg cgaaattcga caaggacact cctgaccgtt
                                                                             180
50
      ggcaaaaaat tgcaagggca gttggtggga aatcaactga agaagtaaag cgacactatg
                                                                             240
      aattgctcct tagggatgtg aatgacattg agtcaggacg ctatccacaa cctagattat
                                                                             300
      gttccgcggc ggcgatagct gctgcgttca cttcaatgtc gatgtcacag aaccgcgctt
                                                                             360
      acgetgacte gegatttege ttteetttet tetettette aceteeggea gaagaatete
                                                                             420
      caaccqatca taagtettet tegaatteta aateggagae taaacetgae teegatgage
                                                                             480
55
      ctaaaggatc tggtttcgat cctgagtcat tggaaagagg tgctaaagct cttcgtgaaa
                                                                             540
                                                                             600
      tcaatagctc tcctcattcc aaacaggtgt ttgatctaat gcgaaagcag gagaagactc
      gattagctga attagcggct gagaaagaac ataatgaagc tattcaagct agcaaggaca
                                                                             660
      ttgaaagaca gcggaaattg gcagaggatc agagaaattt agtgcagcaa caggcccaag
                                                                             720
      cgaaagcgca aaatcttaga tatgaggatg agttggccag gaagagaatg cagacagata
                                                                             780
60
      atgaagctca gagacggcat aatgctgaat tggtttcgat gcaagaagca tcttctatac
                                                                             840
      ggaaagagaa agcaagaatt gccacggaag aacagattca agcacagcag cgcgagactg
                                                                             900
      agaaagagag agcggaactt gagcgagaaa caattcgcgt gaaagctatg gctgaggctg
                                                                             960
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489

```
aaggccgagc tcatgaagct aaactcactg aggagcagaa tagaagaatg cttctagata
                                                                           1020
      agataaatgg tgaaagggag aaatggcttg cagcaatcaa cacgactttc agtcacattg
                                                                           1080
      aaggaggagt aaggacctta ttaactgatc gaagtaaatt gattatgact gttggaggag
                                                                           1140
      ttacggcatt agccgctggg gtttacacaa ctcgagaagg tgctagggtt acctggggtt
                                                                           1200
 5
      atatcaatag aattettggg cagccatcac tgatcagaga atettecatg ggccgattec
                                                                           1260
      catgggcagg ctcagtgtct cagtttaaga acaaacttag cacagctgca ggggcagcag
                                                                           1320
      catctgcaga aggagaaaag cctcttgaaa atgtaattct ccatcgttct ttgaagacga
                                                                           1380
      gaattgagcg tcttgcaaga gccacagcaa ataccaagtc acataaagca ccattccgaa
                                                                           1440
      acatgatgtt ttatgggcct ccaggtaccg gaaaaactat ggtggcgagg gagattgctc
                                                                           1500
10
      ggaagtcggg tcttgattat gctatgatga caggaggaga tgttgctcct cttggtgcac
                                                                           1560
      aggctgttac aaagatccat gaaatatttg attgggctaa gaaatcaaac aaagggttac
                                                                           1620
      tgcttttcat cgatgaagct gatgctttcc tatgcgaacg taacagcact tacatgagtg
                                                                           1680
      aggeteageg eagegetetg aacgegttge tetttegaac eggtgateaa tegegggaca
                                                                           1740
      tagteettgt eetggetaca aacagaeetg gagatetega eagtgeagte aetgaeagga
                                                                           1800
15
     togacgaagt tatogagttt cototocotg gtgaagaaga acgottoaag otootoaago
      tctatctcaa caaqtaccta atgggtgacg acaagaaagg tgagaaagac tcaaacctta
      aatggagcaa cttqttcaag aagaagaagt cacagaagat aaccattgaa ggagacctaa
      ccgaccaagt gattaaagaa gctgcaaaaa agacagaagg cttttctggt cgtgaaatcg
      ctaagcttgt cqccqqtgtt caagctqcqq tatatqqacq acaqqattqt qtcttqqatt
                                                                           2100
20
      cccaactttt tgaagaaatt gtggattata agatcgaaga acatcaccag agaatcagac
                                                                           2160
      ttgcgactga aggtggccaa tcgtttccgt ag
      (2) INFORMATION FOR SEQ ID NO:345:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 703 amino acids
25
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
30
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..703
                (D) OTHER INFORMATION: / Ceres Seq. ID 1974420
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:345:
     Met Ala Ser Ser Met Ser Thr Ser Ser Trp Thr Ala Arg Glu Asp
35
                                          10
      Lys Gln Phe Glu Met Ala Leu Ala Lys Phe Asp Lys Asp Thr Pro Asp
                                      25
      Arg Trp Gln Lys Ile Ala Arg Ala Val Gly Gly Lys Ser Thr Glu Glu
                                  40
40
      Val Lys Arg His Tyr Glu Leu Leu Arg Asp Val Asn Asp Ile Glu
      Ser Gly Arg Tyr Pro Gln Pro Arg Leu Cys Ser Ala Ala Ala Ile Ala
      Ala Ala Phe Thr Ser Met Ser Met Ser Gln Asn Arg Ala Tyr Ala Asp
45
                                          90
      Ser Arg Phe Arg Phe Pro Phe Phe Ser Ser Pro Pro Ala Glu Glu
                                      105
                                                          110
      Ser Pro Thr Asp His Lys Ser Ser Ser Asn Ser Lys Ser Glu Thr Lys
              115
                                  120
50
      Pro Asp Ser Asp Glu Pro Lys Gly Ser Gly Phe Asp Pro Glu Ser Leu
                              135
                                                  140
      Glu Arg Gly Ala Lys Ala Leu Arg Glu Ile Asn Ser Ser Pro His Ser
                                              155
                          150
      Lys Gln Val Phe Asp Leu Met Arg Lys Gln Glu Lys Thr Arg Leu Ala
55
                      165
                                          170
     Glu Leu Ala Ala Glu Lys Glu His Asn Glu Ala Ile Gln Ala Ser Lys
                                      185
      Asp Ile Glu Arg Gln Arg Lys Leu Ala Glu Asp Gln Arg Asn Leu Val
                                 200
                                                      205
60
      Gln Gln Gln Ala Gln Ala Lys Ala Gln Asn Leu Arg Tyr Glu Asp Glu
                              215
                                                  220
```

Leu Ala Arg Lys Arg Met Gln Thr Asp Asn Glu Ala Gln Arg Arg His

										400						
	225					230				490	235					240
			Glu		245					250					255	
5			Arg	260					265					270		
	Thr	Glu	Lys 275	Glu	Arg	Ala	Glu	Leu 280	Glu	Arg	Glu	Thr	Ile 285	Arg	Val	Lys
		290	Ala				295					300				
10	305		Asn			310					315					320
	-	_	Leu		325					330					335	_
15			Thr	340					345					350		
	_		Thr 355					360					365			
20	_	370	Thr	_	_	_	375		_			380				
20	385		Glu			390	_				395		_			400
			Lys		405					410	_				415	
25		_	Glu	420					425					430		•
		_	Ile 435					440					445	_		
2.0	_	450	Pro Met				455			_	_	460		_		_
30	465		Met			470					475		_		_	480
			Ile		485					490					495	
35			Leu	500					505					510		
			515 Tyr			_		520					525			
40		530	Thr				535					540				
- 0	545		Pro	_		550			•		555					560
			Glu	_	565					570					575	
45			Tyr	580				_	585					590		
			595 Ser					600					605			
50		610	Ile				615					620				
- 0	625		Lys			630					635					640
	Val	Ala	Gly	Val	645 Gln	Ala	Ala	Val	Tyr	650 Gly	Arg	Gln	Asp	Cys	655 Val	Leu
55	Asp	Ser	Gln	660 Leu	Phe	Glu	Glu	Ile	665 Val	Asp	Tyr	Lys	Ile	670 Glu	Glu	His
	His	Gln	675 Arg	Ile	Arg	Leu	Ala	680 Thr	Glu	Gly	Gly	Gln	685 Ser	Phe	Pro	
60	(2)	690	) RMA	וא ר ד	F∩Þ	SEO	695 TD 1	ر. ۱0۰۶	16.			700				
	60 (2) INFORMATION FOR SEQ ID NO:346:  (i) SEQUENCE CHARACTERISTICS:															

<sup>(</sup>A) LENGTH: 698 amino acids

491

(B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE:

5

(A) NAME/KEY: peptide

(B) LOCATION: 1..698 (D) OTHER INFORMATION: / Ceres Seq. ID 1974421 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:346: 10 Met Ser Thr Ser Ser Trp Thr Ala Arg Glu Asp Lys Gln Phe Glu Met 10 Ala Leu Ala Lys Phe Asp Lys Asp Thr Pro Asp Arg Trp Gln Lys Ile 25 Ala Arg Ala Val Gly Gly Lys Ser Thr Glu Glu Val Lys Arg His Tyr 15 40 Glu Leu Leu Arg Asp Val Asn Asp Ile Glu Ser Gly Arg Tyr Pro Gln Pro Arq Leu Cys Ser Ala Ala Ala Ile Ala Ala Ala Phe Thr Ser 20 Met Ser Met Ser Gln Asn Arg Ala Tyr Ala Asp Ser Arg Phe Arg Phe Pro Phe Phe Ser Ser Pro Pro Ala Glu Glu Ser Pro Thr Asp His 105 Lys Ser Ser Ser Asn Ser Lys Ser Glu Thr Lys Pro Asp Ser Asp Glu 25 120 Pro Lys Gly Ser Gly Phe Asp Pro Glu Ser Leu Glu Arg Gly Ala Lys 135 Ala Leu Arg Glu Ile Asn Ser Ser Pro His Ser Lys Gln Val Phe Asp 150 155 Leu Met Arg Lys Gln Glu Lys Thr Arg Leu Ala Glu Leu Ala Ala Glu 30 170 175 165 Lys Glu His Asn Glu Ala Ile Gln Ala Ser Lys Asp Ile Glu Arg Gln 185 190 180 Arg Lys Leu Ala Glu Asp Gln Arg Asn Leu Val Gln Gln Gln Ala Gln 35 200 Ala Lys Ala Gln Asn Leu Arg Tyr Glu Asp Glu Leu Ala Arg Lys Arg 215 220 Met Gln Thr Asp Asn Glu Ala Gln Arg Arg His Asn Ala Glu Leu Val 230 235 Ser Met Gln Glu Ala Ser Ser Ile Arg Lys Glu Lys Ala Arg Ile Ala 40 250 Thr Glu Glu Gln Ile Gln Ala Gln Gln Arg Glu Thr Glu Lys Glu Arg 265 Ala Glu Leu Glu Arg Glu Thr Ile Arg Val Lys Ala Met Ala Glu Ala 280 45 Glu Gly Arg Ala His Glu Ala Lys Leu Thr Glu Glu Gln Asn Arg Arg 295 300 Met Leu Leu Asp Lys Ile Asn Gly Glu Arg Glu Lys Trp Leu Ala Ala 310 315 320 50 Ile Asn Thr Thr Phe Ser His Ile Glu Gly Gly Val Arg Thr Leu Leu 325 330 Thr Asp Arg Ser Lys Leu Ile Met Thr Val Gly Gly Val Thr Ala Leu 345 Ala Ala Gly Val Tyr Thr Thr Arg Glu Gly Ala Arg Val Thr Trp Gly 55 360 Tyr Ile Asn Arg Ile Leu Gly Gln Pro Ser Leu Ile Arg Glu Ser Ser 375 380 Met Gly Arg Phe Pro Trp Ala Gly Ser Val Ser Gln Phe Lys Asn Lys 385 390 395 60 Leu Ser Thr Ala Ala Gly Ala Ala Ser Ala Glu Gly Glu Lys Pro 405 410

Leu Glu Asn Val Ile Leu His Arg Ser Leu Lys Thr Arg Ile Glu Arg

				420					425	492				430		
	Leu	Ala	Arg 435		Thr	Ala	Asn	Thr 440		Ser	His	Lys	Ala 445		Phe	Arg
5	Asn	Met 450	Met	Phe	Tyr	Gly	Pro 455	Pro	Gly	Thr	Gly	Lys 460	Thr	Met	Val	Ala
	Arg 465	Glu	Ile	Ala	Arg	Lys 470	Ser	Gly	Leu	Asp	Tyr 475	Ala	Met	Met	Thr	Gly 480
		Asp	Val	Ala	Pro 485	Leu	Gly	Ala	Gln	Ala 490	Val	Thr	Lys	Ile	His 495	Glu
10	Ile	Phe	Asp	Trp 500	Ala	Lys	Lys	Ser	Asn 505	Lys	Gly	Leu	Leu	Leu 510	Phe	Ile
	Asp	Glu	Ala 515	Asp	Ala	Phe	Leu	Cys 520	Glu	Arg	Asn	Ser	Thr 525	Tyr	Met	Ser
15	Glu	Ala 530	Gln	Arg	Ser	Ala	Leu 535	Asn	Ala	Leu	Leu	Phe 540	Arg	Thr	Gly	Asp
	Gln 545	Ser	Arg	Asp	Ile	Val 550	Leu	Val	Leu	Ala	Thr 555	Asn	Arg	Pro	Gly	Asp 560
	Leu	Asp	Ser	Ala	Val 565	Thr	Asp	Arg	Ile	Asp 570	Glu	Val	Ile	Glu	Phe 575	Pro
20	Leu	Pro	Gly	Glu 580	Glu	Glu	Arg	Phe	Lys 585	Leu	Leu	Lys	Leu	Tyr 590	Leu	Asn
	Lys	Tyr	Leu 595	Met	Gly	Asp	Asp	Lys 600	Lys	Gly	Glu	Lys	Asp 605	Ser	Asn	Leu
25	Lys	Trp 610	Ser	Asn	Leu	Phe	Lys 615	Lys	Lys	Lys	Ser	Gln 620	Lys	Ile	Thr	Ile
	Glu 625	Gly	Asp	Leu	Thr	Asp 630	Gln	Val	Ile	Lys	Glu 635	Ala	Ala	Lys	Lys	Thr 640
		_			645					Lys 650					655	
30				660	_	_			665	Val				670		
			675		_			680		Glu	His	His	Gln 685	Arg	Ile	Arg
35		690	Thr		_	_	695			Pro						
	<ul><li>(2) INFORMATION FOR SEQ ID NO:347:</li><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 683 amino acids</li></ul>															
						H: 68 amir			acı	as						
40						DEDNI										
		(ii)	) MOI	LECUI	LE T	OGY: YPE:										
		(ix)	) FEA			KEY:	pept	ide								
45			( I	3) L(	OCAT:	:NOI	1	683							_	
		(xi								Ceres			D 19	74422	2	
					Lys					Thr			Arg	Trp		Lys
50	1 Ile	Ala	Arg		5 Val	Gly	Gly	Lys	Ser 25	10 Thr	Glu	Glu	Val	Lys 30	15 Arg	His
	Tyr	Glu	Leu 35	20 Leu	Leu	Arg	Asp	Val 40		Asp	Ile	Glu	Ser 45	-	Arg	Tyr
55	Pro	Gln 50		Arg	Leu	Cys	Ser 55		Ala	Ala	Ile	Ala 60		Ala	Phe	Thr
	Ser 65		Ser	Met	Ser	Gln 70		Arg	Ala	Tyr	Ala 75		Ser	Arg	Phe	Arg 80
		Pro	Phe	Phe	Ser 85	-	Ser	Pro	Pro	Ala 90	_	Glu	Ser	Pro	Thr 95	
60	His	Lys	Ser	Ser 100		Asn	Ser	Lys	Ser 105	Glu	Thr	Lys	Pro	Asp 110		Asp
	Glu	Pro	Lys		Ser	Gly	Phe	Asp		Glu	Ser	Leu	Glu	Arg	Gly	Ala

								100		495			105			
	Lys	Ala	115 Leu	Arg	Glu	Ile	Asn	120 Ser	Ser	Pro	His	Ser	125 Lys	Gln	Val	Phe
	Asp	130 Leu	Met	Arg	Lys	Gln	135 Glu	Lys	Thr	Arg	Leu	140 Ala	Glu	Leu	Ala	Ala
5	145		Glu	-	-	150				_	155					160
		_			165					170			-		175	_
		_	Lys	180					185					190		
10	Gln	Ala	Lys 195	Ala	Gln	Asn	Leu	Arg 200	Tyr	G1u	Asp	Glu	Leu 205	Ala	Arg	Lys
	Arg	Met 210	Gln	Thr	Asp	Asn	Glu 215	Ala	Gln	Arg	Arg	His 220	Asn	Ala	Glu	Leu
15	Val 225	Ser	Met	Gln	Glu	Ala 230	Ser	Ser	Ile	Arg	Lys 235	Glu	Lys	Ala	Arg	Ile 240
		Thr	Glu	Glu	Gln 245		Gln	Ala	Gln	Gln 250		Glu	Thr	Glu	Lys 255	
	Arg	Ala	Glu	Leu 260		Arg	Glu	Thr	Ile 265		Val	Lys	Ala	Met 270		Glu
20	Ala	Glu	G1y 275		Ala	His	Glu	Ala 280		Leu	Thr	Glu	Glu 285		Asn	Arg
	Arg	Met 290	Leu	Leu	Asp	Lys	Ile 295		Gly	G1u	Arg	Glu 300		Trp	Leu	Ala
25	Ala 305		Asn	Thr	Thr	Phe 310		His	Ile	Glu	Gly 315	Gly	Val	Arg	Thr	Leu 320
		Thr	Asp	Arg	Ser 325	Lys	Leu	Ile	Met	Thr 330		Gly	Gly	Va1	Thr 335	
	Leu	Ala	Ala	Gly 340		Tyr	Thr	Thr	Arg 345		Gly	Ala	Arg	Val 350		Trp
30	Gly	Tyr	Ile 355		Arg	Ile	Leu	Gly 360	Gln	Pro	Ser	Leu	Ile 365		Glu	Ser
	Ser	Met 370	Gly	Arg	Phe	Pro	Trp 375		Gly	Ser	Val	Ser 380		Phe	Lys	Asn
35	Lys 385	Leu	Ser	Thr	Ala	Ala 390	Gly	Ala	Ala	Ala	Ser 395	Ala	Glu	Gly	Glu	Lys 400
	Pro	Leu	Glu	Asn	Val 405	Ile	Leu	His	Arg	Ser 410	Leu	Lys	Thr	Arg	Ile 415	Glu
	Arg	Leu	Ala	Arg 420	Ala	Thr	Ala	Asn	Thr 425	Lys	Ser	His	Lys	Ala 430	Pro	Phe
40	Arg	Asn	Met 435	Met	Phe	Tyr	Gly	Pro 440	Pro	Gly	Thr	Gly	Lys 445	Thr	Met	Val
	Ala	Arg 450	Glu	Ile	Ala	Arg	Lys 455	Ser	Gly	Leu	Asp	Tyr 460	Ala	Met	Met	Thr
45	Gly 465	Gly	Asp	Val	Ala	Pro 470	Leu	Gly	Ala	Gln	Ala 475	Val	Thr	Lys	Ile	His 480
	Glu	Ile	Phe	Asp	Trp 485	Ala	Lys	Lys	Ser	Asn 490	Lys	Gly	Leu	Leu	Leu 495	Phe
	Ile	Asp	Glu	Ala 500	Asp	Ala	Phe	Leu	Cys 505	Glu	Arg	Asn	Ser	Thr 510	Tyr	Met
50	Ser	Glu	Ala 515	Gln	Arg	Ser	Ala	Leu 520	Asn	Ala	Leu	Leu	Phe 525	Arg	Thr	Gly
	Asp	Gln 530	Ser	Arg	Asp	Ile	Val 535	Leu	Val	Leu	Ala	Thr 540	Asn	Arg	Pro	Gly
55	Asp 545		Asp	Ser	Ala	Val 550	Thr	Asp	Arg	Ile	Asp 555	Glu	Val	Ile	Glu	Phe 560
		Leu	Pro	Gly	Glu 565	Glu	Glu	Arg	Phe	Lys 570	Leu	Leu	Lys	Leu	Tyr 575	
	Asn	Lys	Tyr	Leu 580		Gly	Asp	Asp	Lys 585	Lys	Gly	Glu	Lys	Asp 590		Asn
60	Leu	Lys	Trp 595		Asn	Leu	Phe	Lys 600	Lys	Lys	Lys	Ser	Gln 605	Lys	Ile	Thr
	Ile	Glu	Gly	Asp	Leu	Thr	Asp		Val	Ile	Lys	Glu		Ala	Lys	Lys

WO 00/40695 494 615 620 610 Thr Glu Gly Phe Ser Gly Arg Glu Ile Ala Lys Leu Val Ala Gly Val 635 630 Gln Ala Ala Val Tyr Gly Arg Gln Asp Cys Val Leu Asp Ser Gln Leu 5 650 645 Phe Glu Glu Ile Val Asp Tyr Lys Ile Glu Glu His His Gln Arg Ilc 665 Arg Leu Ala Thr Glu Gly Gly Gln Ser Phe Pro 675 (2) INFORMATION FOR SEQ ID NO:348: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 953 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 15 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..953 20 (D) OTHER INFORMATION: / Ceres Seq. ID 1975983 (x1) SEQUENCE DESCRIPTION: SEQ ID NO:348: aaaaaaqata aaacaaaaqc tqaaaqqaqq aaaccaqaqq aaqqaqacqa cgaccaccac cacaacqcat ctcaqctctt ccttaatcca tgqcaqacaa ggqccqcccq ttqccqaaqt 120 ttggaagagc tgagatgcgt tgtggtggtg gcgtcgcctc cttcttcaat ctcatcgcaa 180 25 tcatgatcat atccgagact aatcgccgtg agatctccaa gtacctcttc aaagagggtg 240 ttttgtttgc caaaaaggat ttcaatttac cacaacatcc tttgattgag agtgttccaa 300 atctgcaagt tatcaagttg atgcagagtt tcaaatctaa ggaatatgtg agagagacct 360 ttgcttggat gcattactac tggttcctca caaatgaagg tattgacttt cttaggactt 420 accttaatct cccatctgag attgttcctg ctactctgaa gaagcaacag aagcctcttg 480 30 gtcgaccttt tggaggtggt ggtgaccgtc cccgtggccc tcctcgtggt gatggagaga 540 ggaggtttgg tgacagagat ggataccgtg gaggtcctaa atcaggtgga gagtatggtq 600

acaaggctgg agcacctgct gattaccagc ctggcttcag gggtggagct agtggagcaa 660 ggcaagggtt tggtcgtgga gctggtggtt ttggtggtgg tgctggtcca gctgctggat 720 ctgatctacc ttgaaaagga ctttcttgtt tcttttttggt cttatttaag gttacatagc 780 accttattga gaacgaatgt gtcttttgga actttgtttc tttctcttaa accatttcac 840 aatatgattg ttttttacg ttagaatttt atgaagaaat gttggatcgt tacatgaatc 900 tttttacact gtcctccttt ttaaattctc ctatttagtt tttttcgaag ctt

(2) INFORMATION FOR SEQ ID NO:349:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 214 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- 45 (ix) FEATURE:

35

- (A) NAME/KEY: peptide
- (B) LOCATION: 1..214
- (D) OTHER INFORMATION: / Ceres Seq. ID 1975984
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:349:
- 50 Met Ala Asp Lys Gly Arg Pro Leu Pro Lys Phe Gly Arg Ala Glu Met 10 Arg Cys Gly Gly Val Ala Ser Phe Phe Asn Leu Ile Ala Ile Met 25
- Ile Ile Ser Glu Thr Asn Arg Arg Glu Ile Ser Lys Tyr Leu Phe Lys 55 40
  - Glu Gly Val Leu Phe Ala Lys Lys Asp Phe Asn Leu Pro Gln His Pro 60
  - Leu Ile Glu Ser Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser 75 70
- 60 Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr 85 90 Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg Thr Tyr Leu

495 105 Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys Gln Gln Lys 115 120 Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro Arg Gly Pro 5 135 140 Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp Gly Tyr Arg 150 155 Gly Gly Pro Lys Ser Gly Gly Glu Tyr Gly Asp Lys Ala Gly Ala Pro 170 10 Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly Ala Arg Gln 180 185 Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Ala Gly Pro Ala 195 200 Ala Gly Ser Asp Leu Pro 15 210 (2) INFORMATION FOR SEQ ID NO:350: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 199 amino acids (B) TYPE: amino acid 20 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 25 (B) LOCATION: 1..199 (D) OTHER INFORMATION: / Ceres Seq. ID 1975985 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:350: Met Arg Cys Gly Gly Val Ala Ser Phe Phe Asn Leu Ile Ala Ile 5 10 Met Ile Ile Ser Glu Thr Asn Arg Arg Glu Ile Ser Lys Tyr Leu Phe 30 25 Lys Glu Gly Val Leu Phe Ala Lys Lys Asp Phe Asn Leu Pro Gln His 40 Pro Leu Ile Glu Ser Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln 35 55 Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His 75 7.0 Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg Thr Tyr 90 85 40 Leu Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys Gln Gln 100 105 Lys Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro Arg Gly 120 Pro Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp Gly Tyr 45 135 Arg Gly Gly Pro Lys Ser Gly Gly Glu Tyr Gly Asp Lys Ala Gly Ala 150 155 Pro Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly Ala Arg 170 Gln Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Ala Gly Pro 50 180 185 Ala Ala Gly Ser Asp Leu Pro 195 (2) INFORMATION FOR SEQ ID NO:351: 55 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 183 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 60 (ii) MOLECULE TYPE: peptide (ix) FEATURE:

(A) NAME/KEY: peptide

```
(B) LOCATION: 1..183
                (D) OTHER INFORMATION: / Ceres Seq. ID 1975986
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:351:
      Met Ile Ile Ser Glu Thr Asn Arg Arg Glu Ile Ser Lys Tyr Leu Phe
5
                      5
                                          10
      Lys Glu Gly Val Leu Phe Ala Lys Lys Asp Phe Asn Leu Pro Gln His
                                      25
      Pro Leu Ile Glu Ser Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln
                                  40
10
      Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His
                              55
      Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg Thr Tyr
                          70
                                              75
      Leu Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys Gln Gln
15
      Lys Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro Arg Gly
                                      105
      Pro Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp Gly Tyr
                                  120
20
      Arg Gly Gly Pro Lys Ser Gly Gly Glu Tyr Gly Asp Lys Ala Gly Ala
                              135
      Pro Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly Ala Arg
                         150
                                              155
      Gln Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Ala Gly Pro
25
                                          170
                      165
      Ala Ala Gly Ser Asp Leu Pro
                  180
      (2) INFORMATION FOR SEQ ID NO:352:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 1027 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
35
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1027
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976019
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:352:
40
      atcgaaatta gggtttcgcg ttaggagaag aagttaaagc aaaacacata caaacgcagt
                                                                             60
      caccttetet gtegeeteet tetteaatet categeaate atgateatat eegagaetaa
                                                                             120
      tcqccqtqag atctccaagt acctcttcaa agagggtgtt ttgtttgcca aaaaqgattt
                                                                             180
      caatttacca caacatcctt tgattgagag tgttccaaat ctgcaagtta tcaagttgat
                                                                             240
      gcagagtttc aaatctaagg aatatgtgag agagaccttt gcttggatgc attactactg
                                                                             300
45
      gttcctcaca aatgaaggta ttgactttct taggacttac cttaatctcc catctgagat
                                                                             360
      tgttcctgct actctgaaga agcaacagaa gcctcttggt cgaccttttg gaggtggtgg
                                                                             420
      tgaccgtccc cgtggccctc ctcgtggtga tggagagagg aggtttggtg acagagatgg
                                                                             480
      ataccgtgga ggtcctaaat caggtggaga gtatggtgac aaggctggag cacctgctga
                                                                             540
      ttaccagcct ggcttcaggg gtggagctag tggagcaagg caagggtttg gtcgtggagc
                                                                             600
50
      tggtggtttt ggtggtggtg ctggtccagc tgctggatct gatctacctt gaaaaggaga
                                                                             660
      caatacagct cagccaaagc caaggccttt atcaccttac acaatgtacg cggacatgaa
                                                                             720
      gcctccaaca tcaccacttc catctccagt caccaatcat tagctatgtg agacgagaat
                                                                             780
      tggcctccta aaccctggag cttcttctta attgcaggtt gtaaacaatg agtaagagag
                                                                             840
      tgatggggca attcagtttt gcaggtatga atcagtgagc ttatttgtac aataatacaa
55
      tattcatcaa catqccttat atatqagttc tacttttttt ctcctgttca acqatcaatq
      cagtaaccat attgttaccc tatctcatac atatatgatg acccaaatta attatcatta
                                                                            1020
      (2) INFORMATION FOR SEQ ID NO:353:
           (i) SEQUENCE CHARACTERISTICS:
60
                (A) LENGTH: 183 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
```

```
(D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
 5
               (B) LOCATION: 1..183
               (D) OTHER INFORMATION: / Ceres Seq. ID 1976020
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:353:
     Met Ile Ile Ser Glu Thr Asn Arg Arg Glu Ile Ser Lys Tyr Leu Phe
                                       10
10
     Lys Glu Gly Val Leu Phe Ala Lys Lys Asp Phe Asn Leu Pro Gln His
                                   25
                 20
     Pro Leu Ile Glu Ser Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln
                                 40
     Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His
15
                             55
     Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg Thr Tyr
                                            75
                         70
     Leu Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys Gln Gln
     Lys Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro Arg Gly
20
                                   105
                                                      110
     Pro Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp Gly Tyr
                                120
     Arg Gly Gly Pro Lys Ser Gly Gly Glu Tyr Gly Asp Lys Ala Gly Ala
25
                            135
                                                140
     Pro Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly Ala Arg
                        150 155
     Gln Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Gly Ala Gly Pro
                   165 170
30
     Ala Ala Gly Ser Asp Leu Pro
                 180
      (2) INFORMATION FOR SEQ ID NO:354:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 137 amino acids
35
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
40
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..137
               (D) OTHER INFORMATION: / Ceres Seq. ID 1976021
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:354:
      Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp
45
                                        10
      Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg
                20
                                    25
      Thr Tyr Leu Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys
                                40
50
      Gln Gln Lys Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro
                             55
      Arg Gly Pro Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp
      Gly Tyr Arg Gly Gly Pro Lys Ser Gly Glu Tyr Gly Asp Lys Ala
55
                     85
      Gly Ala Pro Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly
                                    105
      Ala Arg Gln Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Ala
                                120
60
      Gly Pro Ala Ala Gly Ser Asp Leu Pro
                            135
      (2) INFORMATION FOR SEQ ID NO:355:
```

```
(i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 121 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
 5
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..121
10
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976022
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:355:
      Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Asp Phe Leu Arg
                                          10
      Thr Tyr Leu Asn Leu Pro Ser Glu Ile Val Pro Ala Thr Leu Lys Lys
15
                                      25
      Gln Gln Lys Pro Leu Gly Arg Pro Phe Gly Gly Gly Asp Arg Pro
                                  40
      Arg Gly Pro Pro Arg Gly Asp Gly Glu Arg Arg Phe Gly Asp Arg Asp
                              55
                                                  60
20
      Gly Tyr Arg Gly Gly Pro Lys Ser Gly Glu Tyr Gly Asp Lys Ala
                          70
                                              75
      Gly Ala Pro Ala Asp Tyr Gln Pro Gly Phe Arg Gly Gly Ala Ser Gly
                                          90
                     85
      Ala Arg Gln Gly Phe Gly Arg Gly Ala Gly Gly Phe Gly Gly Ala
25
                                     105
                 100
                                                          110
      Gly Pro Ala Ala Gly Ser Asp Leu Pro
              115
      (2) INFORMATION FOR SEQ ID NO:356:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 478 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
35
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..478
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976673
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:356:
40
      aaattcaatc tottocaatt totottotto ttootogoga ogogttaatg goggottogt
                                                                             60
      totcactcac gagetteate teetteatet caecatteaa ateteaaace aaacetacae
                                                                             120
      caccaccaaa totcactott cottotcoaa ctatotocca aaggogaaga aatgatotog
                                                                             180
      ctatcgaatc aatggcggtc gaagaatctt cttcaaccgc ttcttcactt tcctctgagc
                                                                             240
      ttgcttctgt gatatgcccc tcgcttgctt actccaacac gctcttcttc agttctggat
                                                                             300
45
      acaatqtqca aqtqtttqtt gaagataacq aqtcagagga gaggcttqtq aatcqattta
                                                                             360
      qqaqaaqt qatqaqaact qqtgttatac aggaatgtaa qaggaqaaqa tactttqaqa
                                                                             420
      ataaacaaqa tgaqaaqaaa cgtaggactc gtgatgctgc taagcgtaat aaqaaaaq
      (2) INFORMATION FOR SEQ ID NO:357:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 158 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..158
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976674
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:357:
60
      Ile Gln Ser Leu Pro Ile Ser Leu Leu Pro Arg Asp Ala Leu Met
                                          10
      Ala Ala Ser Phe Ser Leu Thr Ser Phe Ile Ser Phe Ile Ser Pro Phe
```

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25
               2.0
     Lys Ser Gln Thr Lys Pro Thr Pro Pro Pro Asn Leu Thr Leu Pro Ser
                    40
     Pro Thr Ile Ser Gln Arg Arg Arg Asn Asp Leu Ala Ile Glu Ser Met
 5
                         55
     Ala Val Glu Ser Ser Ser Thr Ala Ser Ser Leu Ser Ser Glu Leu
     Ala Ser Val Ile Cys Pro Ser Leu Ala Tyr Ser Asn Thr Leu Phe Phe
10
     Ser Ser Gly Tyr Asn Val Gln Val Phe Val Glu Asp Asn Glu Ser Glu
               100
                                105
     Glu Arg Leu Val Asn Arg Phe Arg Arg Glu Val Met Arg Thr Gly Val
                   120
            115
     Ile Gln Glu Cys Lys Arg Arg Tyr Phe Glu Asn Lys Gln Asp Glu
15
      130 135 140
     Lys Lys Arg Arg Thr Arg Asp Ala Ala Lys Arg Asn Lys Lys
          150
     (2) INFORMATION FOR SEQ ID NO:358:
         (i) SEQUENCE CHARACTERISTICS:
20
              (A) LENGTH: 143 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
25
        (ix) FEATURE:
             (A) NAME/KEY: peptide
              (B) LOCATION: 1..143
              (D) OTHER INFORMATION: / Ceres Seq. ID 1976675
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:358:
30
     Met Ala Ala Ser Phe Ser Leu Thr Ser Phe Ile Ser Phe Ile Ser Pro
                 5
                                  10
     Phe Lys Ser Gln Thr Lys Pro Thr Pro Pro Pro Asn Leu Thr Leu Pro
           20
                                25
     Ser Pro Thr Ile Ser Gln Arg Arg Arg Asn Asp Leu Ala Ile Glu Ser
35
                             40
     Met Ala Val Glu Glu Ser Ser Ser Thr Ala Ser Ser Leu Ser Ser Glu
                         55 60
     Leu Ala Ser Val Ile Cys Pro Ser Leu Ala Tyr Ser Asn Thr Leu Phe
                      70
                                      75 80
40
     Phe Ser Ser Gly Tyr Asn Val Gln Val Phe Val Glu Asp Asn Glu Ser
                                    90
                 85
     Glu Glu Arg Leu Val Asn Arg Phe Arg Arg Glu Val Met Arg Thr Gly
               100 105
     Val Ile Gln Glu Cys Lys Arg Arg Arg Tyr Phe Glu Asn Lys Gln Asp
45
           115 120
     Glu Lys Lys Arg Arg Thr Arg Asp Ala Ala Lys Arg Asn Lys Lys
                       135
     (2) INFORMATION FOR SEQ ID NO:359:
          (i) SEQUENCE CHARACTERISTICS:
50
              (A) LENGTH: 95 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
55
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..95
              (D) OTHER INFORMATION: / Ceres Seq. ID 1976676
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:359:
60
     Met Ala Val Glu Glu Ser Ser Ser Thr Ala Ser Ser Leu Ser Ser Glu
                                     10
     Leu Ala Ser Val Ile Cys Pro Ser Leu Ala Tyr Ser Asn Thr Leu Phe
```

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25
                 20
                                                          30
      Phe Ser Ser Gly Tyr Asn Val Gln Val Phe Val Glu Asp Asn Glu Ser
                                  40
      Glu Glu Arg Leu Val Asn Arg Phe Arg Arg Glu Val Met Arg Thr Gly
 5
                              55
      Val Ile Gln Glu Cys Lys Arg Arg Arg Tyr Phe Glu Asn Lys Gln Asp
                                              75
      Glu Lys Lys Arg Arg Thr Arg Asp Ala Ala Lys Arg Asn Lys Lys
                      85
10
      (2) INFORMATION FOR SEQ ID NO:360:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1076 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
15
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1076
20
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025186
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:360:
      aatcatttgc agaagaaaaa taaaaaaaaa aaacagagaa aaggagatat ggaatcaatc
                                                                             60
      ggagtcctta tgatgtgccc catgtcctcc tacctcgaga acgagcttga gaagcgcttc
                                                                             120
      aaccttcttc gcttctggac ttctccggag aaatccgtcc tcctagaaac tcatcggaac
                                                                            180
25
     tocatoogog cogtogttgg gaatgettet geoggegeeg atgeteaget cateagegat
                                                                             240
      ctgcccaacc ttgagattgt atccagcttc agcgtcgggc tcgacaagat cgatttgqqq
                                                                             300
      aaatgcaaag aaaaagggat ccgcgtcacc aacacccccg acgttctcac cgaagacgtc
                                                                             360
      gcagatctcg ccatcgggct tatcctggct ctccttcgac ggctgtgtga gtgcgatcgc
                                                                             420
      tatgtaagga gcggaaaatg gaagcaaggt gaattccaac tcactaccaa gtttagtqga
                                                                             480
30
      aaatccqtqq qqatcattqq tctagqtaga attqqqactq ccatcqcaaa qaqqqctqaa
                                                                             540
      qcctttaqct qcccaatcaa ttactactca agaaccatta agcctgatgt cqcctacaag
                                                                             600
      tattatccqa cqqtqqttqa ccttqctcaa aactcaqaca tcctcqtcqt cqcatqcccq
                                                                             660
      ttgaccgagc agaccagaca cattgtggac cggcaggtca tggatgcatt aggagctaag
                                                                             720
      qqcqtcctca taaacattqq ccqtqqacca catqttqatq aqcaaqaqct tattaaaqct
                                                                             780
35
      ctaacaqaaq qccqcctaqq tggggctgcc cttgatgtgt ttgagcagqa gcctcacqtq
      cccqaqqaqc tctttqqcct tgagaatgta gttctcctcc ctcacgttqq gaqtqqcact
                                                                             900
      qtqqaaacac qqaatqccat qqccqatctt qtcqtqqgta acttqqaaqc qcacttttct
                                                                            960
      qqqaaatcac ttctqactcc qqtcqtctqa qtcatqtqac acatttqqqa tttqqtttca
                                                                            1020
      cttcatttga aaggttttta tgttaacgaa ctcatgaaaa gggtaatttc tcttcc
      (2) INFORMATION FOR SEQ ID NO:361:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 329 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..329
50
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025187
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:361:
      Asn His Leu Gln Lys Lys Asn Lys Lys Lys Gln Arg Lys Gly Asp
                                          10
      Met Glu Ser Ile Gly Val Leu Met Met Cys Pro Met Ser Ser Tyr Leu
55
                                      25
      Glu Asn Glu Leu Glu Lys Arg Phe Asn Leu Leu Arg Phe Trp Thr Ser
                                  40
      Pro Glu Lys Ser Val Leu Leu Glu Thr His Arg Asn Ser Ile Arg Ala
                              55
                                                  60
60
      Val Val Gly Asn Ala Ser Ala Gly Ala Asp Ala Gln Leu Ile Ser Asp
                                               75
      Leu Pro Asn Leu Glu Ile Val Ser Ser Phe Ser Val Gly Leu Asp Lys
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501 90 85 Ile Asp Leu Gly Lys Cys Lys Glu Lys Gly Ile Arg Val Thr Asn Thr 100 105 110 Pro Asp Val Leu Thr Glu Asp Val Ala Asp Leu Ala Ile Gly Leu Ile 5 115 120 Leu Ala Leu Leu Arg Arg Leu Cys Glu Cys Asp Arg Tyr Val Arg Ser 135 Gly Lys Trp Lys Gln Gly Glu Phe Gln Leu Thr Thr Lys Phe Ser Gly 150 155 160 Lys Ser Val Gly Ile Ile Gly Leu Gly Arg Ile Gly Thr Ala Ile Ala 10 165 170 175 Lys Arg Ala Glu Ala Phe Ser Cys Pro Ile Asn Tyr Tyr Ser Arg Thr 185 Ile Lys Pro Asp Val Ala Tyr Lys Tyr Tyr Pro Thr Val Val Asp Leu 15 200 Ala Gln Asn Ser Asp Ile Leu Val Val Ala Cys Pro Leu Thr Glu Gln 215 Thr Arg His Ile Val Asp Arg Gln Val Met Asp Ala Leu Gly Ala Lys 230 235 20 Gly Val Leu Ile Asn Ile Gly Arg Gly Pro His Val Asp Glu Gln Glu 245 250 255 Leu Ile Lys Ala Leu Thr Glu Gly Arg Leu Gly Gly Ala Ala Leu Asp 260 265 Val Phe Glu Gln Glu Pro His Val Pro Glu Glu Leu Phe Gly Leu Glu 25 275 280 Asn Val Val Leu Leu Pro His Val Gly Ser Gly Thr Val Glu Thr Arg 290 295 300 Asn Ala Met Ala Asp Leu Val Val Gly Asn Leu Glu Ala His Phe Ser 305 310 315 30 Gly Lys Ser Leu Leu Thr Pro Val Val 325 (2) INFORMATION FOR SEQ ID NO:362: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 313 amino acids 35 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 40 (A) NAME/KEY: peptide (B) LOCATION: 1..313 (D) OTHER INFORMATION: / Ceres Seq. ID 2025188 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:362: Met Glu Ser Ile Gly Val Leu Met Met Cys Pro Met Ser Ser Tyr Leu 10 45 Glu Asn Glu Leu Glu Lys Arg Phe Asn Leu Leu Arg Phe Trp Thr Ser 25 Pro Glu Lys Ser Val Leu Leu Glu Thr His Arg Asn Ser Ile Arg Ala 40 50 Val Val Gly Asn Ala Ser Ala Gly Ala Asp Ala Gln Leu Ile Ser Asp 55 Leu Pro Asn Leu Glu Ile Val Ser Ser Phe Ser Val Gly Leu Asp Lys 70 75 Ile Asp Leu Gly Lys Cys Lys Glu Lys Gly Ile Arg Val Thr Asn Thr 55 90 Pro Asp Val Leu Thr Glu Asp Val Ala Asp Leu Ala Ile Gly Leu Ile 105 Leu Ala Leu Leu Arg Arg Leu Cys Glu Cys Asp Arg Tyr Val Arg Ser

120

135

60

Gly Lys Trp Lys Gln Gly Glu Phe Gln Leu Thr Thr Lys Phe Ser Gly

Lys Ser Val Gly Ile Ile Gly Leu Gly Arg Ile Gly Thr Ala Ile Ala

502 150 155 Lys Arg Ala Glu Ala Phe Ser Cys Pro Ile Asn Tyr Tyr Ser Arg Thr 170 165 Ile Lys Pro Asp Val Ala Tyr Lys Tyr Tyr Pro Thr Val Val Asp Leu 185 5 180 Ala Gln Asn Ser Asp Ile Leu Val Val Ala Cys Pro Leu Thr Glu Gln 200 Thr Arg His Ile Val Asp Arg Gln Val Met Asp Ala Leu Gly Ala Lys 215 10 Gly Val Leu Ile Asn Ile Gly Arg Gly Pro His Val Asp Glu Gln Glu 235 230 Leu Ile Lys Ala Leu Thr Glu Gly Arg Leu Gly Gly Ala Ala Leu Asp 245 250 Val Phe Glu Glu Glu Pro His Val Pro Glu Glu Leu Phe Gly Leu Glu 15 265 Asn Val Val Leu Pro His Val Gly Ser Gly Thr Val Glu Thr Arg 280 Asn Ala Met Ala Asp Leu Val Val Gly Asn Leu Glu Ala His Phe Ser 290 295 20 Gly Lys Ser Leu Leu Thr Pro Val Val 310 (2) INFORMATION FOR SEQ ID NO:363: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 306 amino acids 25 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 30 (A) NAME/KEY: peptide (B) LOCATION: 1..306 (D) OTHER INFORMATION: / Ceres Seq. ID 2025189 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:363: Met Met Cys Pro Met Ser Ser Tyr Leu Glu Asn Glu Leu Glu Lys Arg 35 10 Phe Asn Leu Leu Arg Phe Trp Thr Ser Pro Glu Lys Ser Val Leu Leu 20 25 Glu Thr His Arg Asn Ser Ile Arg Ala Val Val Gly Asn Ala Ser Ala 4.0 40 Gly Ala Asp Ala Gln Leu Ile Ser Asp Leu Pro Asn Leu Glu Ile Val 55 Ser Ser Phe Ser Val Gly Leu Asp Lys Ile Asp Leu Gly Lys Cys Lys 70 75 Glu Lys Gly Ile Arg Val Thr Asn Thr Pro Asp Val Leu Thr Glu Asp 45 90 Val Ala Asp Leu Ala Ile Gly Leu Ile Leu Ala Leu Leu Arg Arg Leu 100 105 Cys Glu Cys Asp Arg Tyr Val Arg Ser Gly Lys Trp Lys Gln Gly Glu 120 50 Phe Gln Leu Thr Thr Lys Phe Ser Gly Lys Ser Val Gly Ile Ile Gly 135 Leu Gly Arg Ile Gly Thr Ala Ile Ala Lys Arg Ala Glu Ala Phe Ser 150 155 Cys Pro Ile Asn Tyr Tyr Ser Arg Thr Ile Lys Pro Asp Val Ala Tyr 170 55 165 Lys Tyr Tyr Pro Thr Val Val Asp Leu Ala Gln Asn Ser Asp Ile Leu 185 180 190 Val Val Ala Cys Pro Leu Thr Glu Gln Thr Arg His Ile Val Asp Arg 200 205 60 Gln Val Met Asp Ala Leu Gly Ala Lys Gly Val Leu Ile Asn Ile Gly 220 215

Arg Gly Pro His Val Asp Glu Glu Leu Ile Lys Ala Leu Thr Glu

503 230 235 Gly Arg Leu Gly Gly Ala Ala Leu Asp Val Phe Glu Gln Glu Pro His 250 245 Val Pro Glu Glu Leu Phe Gly Leu Glu Asn Val Val Leu Leu Pro His 5 265 260 Val Gly Ser Gly Thr Val Glu Thr Arg Asn Ala Met Ala Asp Leu Val 280 Val Gly Asn Leu Glu Ala His Phe Ser Gly Lys Ser Leu Leu Thr Pro 10 Val Val 305 (2) INFORMATION FOR SEQ ID NO:364: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 555 base pairs 15 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 20 (A) NAME/KEY: -(B) LOCATION: 1..555 (D) OTHER INFORMATION: / Ceres Seq. ID 2025372 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:364: aaaactcact ttcacttgca caaagataag gaaaccatgt ctgtgtcagc gatctttggt 60 25 accggaatcg tcaccgttgc tgcttctccg gttctccgcc aatttcaagt tccaaaattg 120 ggtaatggag gaggattagg gatggtgata gagtgttcgt cgaggccaca gaagaaatcg 180 acggctcatc acaggaagac gaggccgaag aagactcagc cttgggacat taagagaaag 240 cctactgttt acgctcctct tcctcctctt ccggcggaat ggagtccgtt cactctcgct 300 tctgacgacg gtggtgctgc cactgctgcg ggagatttgg tttcaggcgc tgcctagttg 360 30 gtatgagtta tetgetgatt tggttgtaat etagtttgag aacttitgtt gttgtttact 420 480 ggaccaaatt tgcgatttat aaagcttcaa cctttctcct ctgtttttga gtttaaaggt 540 ctcaacttta tagtt (2) INFORMATION FOR SEQ ID NO:365: 35 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 118 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 40 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..118 (D) OTHER INFORMATION: / Ceres Seq. ID 2025373 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:365: Lys Thr His Phe His Leu His Lys Asp Lys Glu Thr Met Ser Val Ser 10 Ala Ile Phe Gly Thr Gly Ile Val Thr Val Ala Ala Ser Pro Val Leu 25 50 Arg Gln Phe Gln Val Pro Lys Leu Gly Asn Gly Gly Gly Leu Gly Met 40 Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser Thr Ala His His Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp Ile Lys Arg Lys 55 75 70 Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala Glu Trp Ser Pro 90 Phe Thr Leu Ala Ser Asp Gly Gly Ala Ala Thr Ala Ala Gly Asp 100 105 60 Leu Val Ser Gly Ala Ala 115 (2) INFORMATION FOR SEQ ID NO: 366:

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(i) SEOUENCE CHARACTERISTICS:
                (A) LENGTH: 106 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..106
10
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025374
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:366:
      Met Ser Val Ser Ala Ile Phe Gly Thr Gly Ile Val Thr Val Ala Ala
                                           10
      Ser Pro Val Leu Arg Gln Phe Gln Val Pro Lys Leu Gly Asn Gly Gly
15
                                      25
      Gly Leu Gly Met Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser
                                  40
      Thr Ala His His Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp
                              55
20
      Ile Lys Arg Lys Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala
                                               75
                          70
      Glu Trp Ser Pro Phe Thr Leu Ala Ser Asp Asp Gly Gly Ala Ala Thr
                                          90
      Ala Ala Gly Asp Leu Val Ser Gly Ala Ala
25
                  100
      (2) INFORMATION FOR SEQ ID NO:367:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 71 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
35
                (B) LOCATION: 1..71
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025375
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:367:
      Met Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser Thr Ala His
                                           10
40
      His Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp Ile Lys Arg
                                       25
      Lys Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala Glu Trp Ser
                                  4.0
      Pro Phe Thr Leu Ala Ser Asp Asp Gly Gly Ala Ala Thr Ala Ala Gly
45
                              5.5
      Asp Leu Val Ser Gly Ala Ala
                          70
      (2) INFORMATION FOR SEQ ID NO:368:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 631 base pairs
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: DNA (genomic)
55
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..631
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025471
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:368:
      atgtcgcgaa agttcctaaa accgccacgt gtcagtcact tatccaacag taatccaatt
60
                                                                              60
      attegteaag ceteeggttt attacegtet aaacegaaaa gaaaaaacte tetteggaat
                                                                              120
      ctgaaaagcc ataaatcagc agaaaaaaat cttaaggatc ggaaatttta taatcggatg
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505

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aatagagtga ttagtcagtt ttccggtaag ttgatgaagg agaagagtgt taccggtqta
     ageggeggag atacqaettt aagatetgee gteaaagaat eggtetegte teeacaqagt
                                                                            300
     gcgtcgtcgt cgtcgtcggt gaggaggctt aaaggagatc tggaatcgag tcgatttggt
                                                                            360
     gcggcggcga gtgagaggct gaggcaagcg gaggaatctt tgaggacggt gatgtttctq
                                                                            420
 5
     agctgttggg gatcttgtta gatctggttg agaaaataat agatgagaaa acgaaaccaa
                                                                            480
     aacgtttctg tttgtgtttt tgagttttgg atttttgttt tctctgtgaa tagtttttt
                                                                            540
     ttttttttttc gttttctttg tactttgtgt tttatgtaaa tcatgtataa aatgaaaagt
                                                                            600
     aaatcaatga agataatggt ttgtcttcac c
      (2) INFORMATION FOR SEQ ID NO:369:
10
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 146 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
15
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..146
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025472
20
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:369:
     Met Ser Arg Lys Phe Leu Lys Pro Pro Arg Val Ser His Leu Ser Asn
                                          10
      Ser Asn Pro Ile Ile Arg Gln Ala Ser Gly Leu Leu Pro Ser Lys Pro
                                      25
                 20
25
      Lys Arg Lys Asn Ser Leu Arg Asn Leu Lys Ser His Lys Ser Ala Glu
                                  40
      Lys Asn Leu Lys Asp Arg Lys Phe Tyr Asn Arg Met Asn Arg Val Ile
      Ser Gln Phe Ser Gly Lys Leu Met Lys Glu Lys Ser Val Thr Gly Val
30
                          70
      Ser Gly Gly Asp Thr Thr Leu Arg Ser Ala Val Lys Glu Ser Val Ser
                                          90
      Ser Pro Gln Ser Ala Ser Ser Ser Ser Ser Val Arg Arg Leu Lys Gly
                 100
                                     105
35
      Asp Leu Glu Ser Ser Arg Phe Gly Ala Ala Ser Glu Arg Leu Arg
                              120
                                                     125
      Gln Ala Glu Glu Ser Leu Arg Thr Val Met Phe Leu Ser Cys Trp Gly
                             135
          130
      Ser Cys
40
      145
      (2) INFORMATION FOR SEQ ID NO: 370:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 87 amino acids
                (B) TYPE: amino acid
45
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
50
                (B) LOCATION: 1..87
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025473
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:370:
      Met Asn Arg Val Ile Ser Gln Phe Ser Gly Lys Leu Met Lys Glu Lys
                                          10
55
      Ser Val Thr Gly Val Ser Gly Gly Asp Thr Thr Leu Arg Ser Ala Val
                  20
      Lys Glu Ser Val Ser Ser Pro Gln Ser Ala Ser Ser Ser Ser Val
                                  40
      Arg Arg Leu Lys Gly Asp Leu Glu Ser Ser Arg Phe Gly Ala Ala Ala
60
                              55
      Ser Glu Arg Leu Arg Gln Ala Glu Glu Ser Leu Arg Thr Val Met Phe
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506

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Leu Ser Cys Trp Gly Ser Cys
                      85
      (2) INFORMATION FOR SEQ ID NO: 371:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 75 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
10
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025474
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:371:
15
     Met Lys Glu Lys Ser Val Thr Gly Val Ser Gly Gly Asp Thr Thr Leu
                                           10
      Arg Ser Ala Val Lys Glu Ser Val Ser Ser Pro Gln Ser Ala Ser Ser
      Ser Ser Ser Val Arg Arg Leu Lys Gly Asp Leu Glu Ser Ser Arg Phe
20
                                  40
      Gly Ala Ala Ser Glu Arg Leu Arg Gln Ala Glu Glu Ser Leu Arg
                              55
                                                   60
      Thr Val Met Phe Leu Ser Cys Trp Gly Ser Cys
                          70
25
      (2) INFORMATION FOR SEQ ID NO:372:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 2029 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..2029
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025475
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:372:
      atgggtaaga gttggttttc agctgtgaag aaagcattaa gcccagaacc aaaacaaaag
                                                                              60
      aaaqaqcaqa aqccacataa qtccaaqaaa tqqtttqqta aatccaaqaa qctaqatqtt
                                                                             120
      actaattetg gtgcagcata ttetectegt actgtcaaag acgegaaact aaaggagatt
                                                                             180
40
      gaggagcaac agagcagaca tgcttactct gtggctattg caactgctgc agctgcagag
                                                                             240
      gcagccgttg cagctgctca agctgctgct gaagttgttc gtctctctgc attatcacqq
                                                                             300
      ttccccqqqa aatcaatgga agagatcgcc gctatcaaga ttcagacagc atttagagga
                                                                             360
      tatatggcaa qaagagcatt gcgtgcgttg agaggtcttg tgaggctaaa atctttagtc
                                                                             420
      caggggaaat gtgtgagacg tcaagccaca tctacattgc aaagcatgca aacactagct
                                                                             480
45
      agagtacaat atcagattcg tgagagaagg ctccgattgt ctgaggataa acaggcttta
                                                                             540
      acacgacage tecaacaaaa acacaataaa gaetttgata agaetggaga aaattggaat
                                                                             600
      gatagtacat tgtcgcggga gaaagttgaa gcaaacatgt tgaacaagca agtagcaaca
                                                                             660
                                                                             720
      atgagaagag aaaaagcgct tgcatatgca ttcagtcacc agaatacatg gaaaaactca
      actaaaatgg gttctcaaac attcatggac cctaacaatc cgcattgggg ttggagttgg
                                                                             780
50
      ctagaacgtt ggatggctgc tcgaccaaac gaaaaccact cactcacacc agataatgct
                                                                             840
      gaaaaagact cttctgctag gagtgtagca agccgtgcca tgtctgagat gattccacga
                                                                             900
      ggcaaaaacc tttcaccaag aggaaagaca ccaaacagtc gaagagggtc aagcccgaga
                                                                             960
      gtgaggcaag tcccaagtga agactcaaac agcattgtga gtttccaatc agaacaacct
                                                                            1020
      tgcaatcgta ggcatagcac ttgtggatca attccatcaa ctagagacga tgaaagcttc
                                                                            1080
55
      accagtagtt teteteagte agtteeagge tacatggeae etacacaage egecaaagea
                                                                            1140
      agagetegat teteaaacet tagteeteta ageteagaga agacagegaa aaaacegett
                                                                            1200
      tccttctcgg gatctcctaa gactgtaaga cggttttcag tagggattct agggctttcc
                                                                            1260
      ttcattggtc agatcagacg acgttttaca tcttcttctt cttcctcttc gatattcgtc
                                                                            1320
      aqtqtqtqta ttttqqqqaa aactttgtga qcaaaqagcg agaaaatgag cqqaccqqta
                                                                            1380
60
      agaaaatcgc ggatgtggct ttcaaagctt cgaggactat cgattgggat ggtatggcta
                                                                            1440
      aggtccttgt cacagatgag gctcgtagag agttctctaa ccttcgtcgt gctttcgatg
                                                                            1500
```

aggttaacac acagctccag accaaattta gtcaggaacc tgaacccata gattgggatt

507

```
actataggaa gggtattgga gctggcattg ttgacaagta caaggaagct tatgacagca
      ttgagattcc aaagtacgtt gacaaagtta ctcctgaata caagccaaag tttgatgctt
                                                                           1680
      tqttqqtqqa actqaaaqaa gcagaacaga aatcqctcaa ggagtctgaa cggttggaga
                                                                           1740
      aagaaattgc tgatgtccaa gagatcagca aaaagctcag caccatgact gcagatgagt
 5
      actttgagaa gcacccggaa ctcaaaaaga agtttgatga cgaaatccgt aatgacaact
                                                                           1860
      ggggatactg atcatgtttc tccatctccg gcttggaaag aaaactctct ttctctttct
                                                                           1920
      ctgttctctt actgtgattt tgtgagccaa tcataacaat aataagtaca ccattcactt
      aagcagtgtt gagatcttca ttccaaggaa gataaacgca tttggtttt
      (2) INFORMATION FOR SEQ ID NO:373:
10
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 449 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
15
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..449
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025476
20
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:373:
      Met Gly Lys Ser Trp Phe Ser Ala Val Lys Lys Ala Leu Ser Pro Glu
                                          10
      Pro Lys Gln Lys Lys Glu Gln Lys Pro His Lys Ser Lys Lys Trp Phe
                                      25
25
      Gly Lys Ser Lys Lys Leu Asp Val Thr Asn Ser Gly Ala Ala Tyr Ser
                                  40
      Pro Arg Thr Val Lys Asp Ala Lys Leu Lys Glu Ile Glu Glu Gln Gln
      Ser Arg His Ala Tyr Ser Val Ala Ile Ala Thr Ala Ala Ala Glu
30
                          70
                                              75
      Ala Ala Val Ala Ala Ala Gln Ala Ala Glu Val Val Arg Leu Ser
                      85
                                          90
      Ala Leu Ser Arg Phe Pro Gly Lys Ser Met Glu Glu Ile Ala Ala Ile
                  100
                                      105
                                                          110
35
      Lys Ile Gln Thr Ala Phe Arg Gly Tyr Met Ala Arg Arg Ala Leu Arg
                                  120
                                                      125
      Ala Leu Arg Gly Leu Val Arg Leu Lys Ser Leu Val Gln Gly Lys Cys
                              135
                                                  140
      Val Arg Arg Gln Ala Thr Ser Thr Leu Gln Ser Met Gln Thr Leu Ala
40
                          150
                                             155
      Arg Val Gln Tyr Gln Ile Arg Glu Arg Arg Leu Arg Leu Ser Glu Asp
                                          170
                      165
      Lys Gln Ala Leu Thr Arg Gln Leu Gln Gln Lys His Asn Lys Asp Phe
                  180
                                      185
                                                         190
45
      Asp Lys Thr Gly Glu Asn Trp Asn Asp Ser Thr Leu Ser Arg Glu Lys
                                  200
                                                      205
      Val Glu Ala Asn Met Leu Asn Lys Gln Val Ala Thr Met Arg Arg Glu
                              215
                                                  220
      Lys Ala Leu Ala Tyr Ala Phe Ser His Gln Asn Thr Trp Lys Asn Ser
50
                          230
                                              235
      Thr Lys Met Gly Ser Gln Thr Phe Met Asp Pro Asn Asn Pro His Trp
                      245
                                          250
      Gly Trp Ser Trp Leu Glu Arg Trp Met Ala Ala Arg Pro Asn Glu Asn
                                      265
                  260
55
      His Ser Leu Thr Pro Asp Asn Ala Glu Lys Asp Ser Ser Ala Arg Ser
                                  280
      Val Ala Ser Arg Ala Met Ser Glu Met Ile Pro Arg Gly Lys Asn Leu
                              295
      Ser Pro Arg Gly Lys Thr Pro Asn Ser Arg Arg Gly Ser Ser Pro Arg
60
                          310
                                              315
      Val Arg Gln Val Pro Ser Glu Asp Ser Asn Ser Ile Val Ser Phe Gln
```

330

508 Ser Glu Gln Pro Cys Asn Arg Arg His Ser Thr Cys Gly Ser Ile Pro 345 350 Ser Thr Arg Asp Asp Glu Ser Phe Thr Ser Ser Phe Ser Gln Ser Val 355 360 365 5 Pro Gly Tyr Met Ala Pro Thr Gln Ala Ala Lys Ala Arg Ala Arg Phe 370 375 380 Ser Asn Leu Ser Pro Leu Ser Ser Glu Lys Thr Ala Lys Lys Arg Leu 390 395 Ser Phe Ser Gly Ser Pro Lys Thr Val Arg Arg Phe Ser Val Gly Ile 10 410 415 Leu Gly Leu Ser Phe Ile Gly Gln Ile Arg Arg Arg Phe Thr Ser Ser 420 425 Ser Ser Ser Ser Ile Phe Val Ser Val Cys Ile Leu Gly Lys Thr 440 15 Leu (2) INFORMATION FOR SEQ ID NO:374: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 344 amino acids 20 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 25 (A) NAME/KEY: peptide (B) LOCATION: 1..344 (D) OTHER INFORMATION: / Ceres Seq. ID 2025477 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:374: Met Glu Glu Ile Ala Ala Ile Lys Ile Gln Thr Ala Phe Arg Gly Tyr 10 30 Met Ala Arg Arg Ala Leu Arg Ala Leu Arg Gly Leu Val Arg Leu Lys 20 25 Ser Leu Val Gln Gly Lys Cys Val Arg Arg Gln Ala Thr Ser Thr Leu 40 35 Gln Ser Met Gln Thr Leu Ala Arg Val Gln Tyr Gln Ile Arg Glu Arg 55 Arg Leu Arg Leu Ser Glu Asp Lys Gln Ala Leu Thr Arg Gln Leu Gln 75 70 Gln Lys His Asn Lys Asp Phe Asp Lys Thr Gly Glu Asn Trp Asn Asp 40 90 85 Ser Thr Leu Ser Arg Glu Lys Val Glu Ala Asn Met Leu Asn Lys Gln 105 Val Ala Thr Met Arg Arg Glu Lys Ala Leu Ala Tyr Ala Phe Ser His 120 45 Gln Asn Thr Trp Lys Asn Ser Thr Lys Met Gly Ser Gln Thr Phe Met 135 Asp Pro Asn Asn Pro His Trp Gly Trp Ser Trp Leu Glu Arg Trp Met 155 Ala Ala Arg Pro Asn Glu Asn His Ser Leu Thr Pro Asp Asn Ala Glu 50 170 175 Lys Asp Ser Ser Ala Arg Ser Val Ala Ser Arg Ala Met Ser Glu Met 185 Ile Pro Arg Gly Lys Asn Leu Ser Pro Arg Gly Lys Thr Pro Asn Ser 200 55 Arg Arg Gly Ser Ser Pro Arg Val Arg Gln Val Pro Ser Glu Asp Ser 215 220 Asn Ser Ile Val Ser Phe Gln Ser Glu Gln Pro Cys Asn Arg Arg His 230 235 240 Ser Thr Cys Gly Ser Ile Pro Ser Thr Arg Asp Asp Glu Ser Phe Thr 60 245 250 255 Ser Ser Phe Ser Gln Ser Val Pro Gly Tyr Met Ala Pro Thr Gln Ala 265 260 270

509 Ala Lys Ala Arg Ala Arg Phe Ser Asn Leu Ser Pro Leu Ser Ser Glu 280 285 Lys Thr Ala Lys Lys Arg Leu Ser Phe Ser Gly Ser Pro Lys Thr Val 290 295 300 5 Arg Arg Phe Ser Val Gly Ile Leu Gly Leu Ser Phe Ile Gly Gln Ile 310 315 Arg Arg Arg Phe Thr Ser Ser Ser Ser Ser Ser Ile Phe Val Ser 330 325 Val Cys Ile Leu Gly Lys Thr Leu 10 340 (2) INFORMATION FOR SEQ ID NO:375: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 328 amino acids (B) TYPE: amino acid 15 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 20 (B) LOCATION: 1..328 (D) OTHER INFORMATION: / Ceres Seq. ID 2025478 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:375: Met Ala Arg Arg Ala Leu Arg Ala Leu Arg Gly Leu Val Arg Leu Lys 10 Ser Leu Val Gln Gly Lys Cys Val Arg Arg Gln Ala Thr Ser Thr Leu 25 25 20 Gln Ser Met Gln Thr Leu Ala Arg Val Gln Tyr Gln Ile Arg Glu Arg 40 Arg Leu Arg Leu Ser Glu Asp Lys Gln Ala Leu Thr Arg Gln Leu Gln 30 55 Gln Lys His Asn Lys Asp Phe Asp Lys Thr Gly Glu Asn Trp Asn Asp 75 80 70 Ser Thr Leu Ser Arg Glu Lys Val Glu Ala Asn Met Leu Asn Lys Gln 90 35 Val Ala Thr Met Arg Arg Glu Lys Ala Leu Ala Tyr Ala Phe Ser His 105 Gln Asn Thr Trp Lys Asn Ser Thr Lys Met Gly Ser Gln Thr Phe Met 120 Asp Pro Asn Asn Pro His Trp Gly Trp Ser Trp Leu Glu Arg Trp Met 40 135 Ala Ala Arg Pro Asn Glu Asn His Ser Leu Thr Pro Asp Asn Ala Glu 150 155 Lys Asp Ser Ser Ala Arg Ser Val Ala Ser Arg Ala Met Ser Glu Met 165 170 175 45 Ile Pro Arg Gly Lys Asn Leu Ser Pro Arg Gly Lys Thr Pro Asn Ser 185 180 Arg Arg Gly Ser Ser Pro Arg Val Arg Gln Val Pro Ser Glu Asp Ser 200 Asn Ser Ile Val Ser Phe Gln Ser Glu Gln Pro Cys Asn Arg Arg His 50 215 Ser Thr Cys Gly Ser Ile Pro Ser Thr Arg Asp Asp Glu Ser Phe Thr 230 235 Ser Ser Phe Ser Gln Ser Val Pro Gly Tyr Met Ala Pro Thr Gln Ala 250 245 55 Ala Lys Ala Arg Ala Arg Phe Ser Asn Leu Ser Pro Leu Ser Ser Glu 260 265 Lys Thr Ala Lys Lys Arg Leu Ser Phe Ser Gly Ser Pro Lys Thr Val 280 285 Arg Arg Phe Ser Val Gly Ile Leu Gly Leu Ser Phe Ile Gly Gln Ile 290 295 300 60 Arg Arg Phe Thr Ser Ser Ser Ser Ser Ser Ile Phe Val Ser

310

305

315

510

```
Val Cys Ile Leu Gly Lys Thr Leu
                      325
      (2) INFORMATION FOR SEQ ID NO:376:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 678 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
10
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..678
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025524
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:376:
15
     gattttttag ggtttcaagt gaaaagagta atagcgcggc ggagccatgg ttctcaagac
                                                                             60
      tgagctttgc cgattcagtg gccagaaaat ttaccctggt agagggatca gatttatccg
                                                                             120
     atcggactct caggtgtttt tgtttctcaa ctccaaatgt aagaggtatt tccacaacaa
                                                                             180
     gttgaagcca tctaagcttt gctggactgc tatgtaccga aagcagcaca agaaggacgc
                                                                             240
      agcacaagag gctgtgaaga gaaggagacg tgcaactaag aagccttact caaggtcgat
                                                                             300
20
     tgtcggtgct actttggagg ttattcagaa gaagcgagca gagaagcctg aagttcgtga
                                                                             360
      tgccgctaga gaagctgccc tacgtgagat caaggagaga atcaagaaga ccaaggacga
                                                                            420
      gaagaaggca aagaaggtcg agtatgcatc aaagcaacag aagtcacaag tgaagggaaa
                                                                            480
      tatccccaag agtgctgcac ccaaggctgc taagatgggt ggtggtggag gcagacgttg
                                                                            540
      aatggageta tagagtagee caetettete tetteaetta tetttette ttgttttgae
                                                                             600
25
      attgttttgt tttgtcagcc attttttagt tttgcaccag atctaatata ttcagtttat
                                                                             660
      gaaaactttt tgtttggc
      (2) INFORMATION FOR SEQ ID NO:377:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 164 amino acids
30
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..164
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025525
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:377:
      Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr
40
                                          10
      Pro Gly Arg Gly Ile Arg Phe Ile Arg Ser Asp Ser Gln Val Phe Leu
                                      25
      Phe Leu Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Lys Leu Lys Pro
                                  40
45
      Ser Lys Leu Cys Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp
                              55
      Ala Ala Gln Glu Ala Val Lys Arg Arg Arg Ala Thr Lys Lys Pro
                          70
                                               75
      Tyr Ser Arg Ser Ile Val Gly Ala Thr Leu Glu Val Ile Gln Lys Lys
50
                                          90
      Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu
                                      105
                  100
      Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala
                                  120
              115
55
      Lys Lys Val Glu Tyr Ala Ser Lys Gln Gln Lys Ser Gln Val Lys Gly
                              135
                                                  140
      Asn Ile Pro Lys Ser Ala Ala Pro Lys Ala Ala Lys Met Gly Gly Gly
                          150
                                               155
      Gly Gly Arg Arg
60
```

(2) INFORMATION FOR SEQ ID NO:378:

(i) SEQUENCE CHARACTERISTICS:

```
(A) LENGTH: 109 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..109
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025526
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:378:
      Met Tyr Arg Lys Gln His Lys Lys Asp Ala Ala Gln Glu Ala Val Lys
                                          10
      Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
                  20
                                      25
15
      Ala Thr Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val
                                  40
      Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
      Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Val Glu Tyr Ala Ser
20
                          70
                                              75
      Lys Gln Gln Lys Ser Gln Val Lys Gly Asn Ile Pro Lys Ser Ala Ala
                      85
      Pro Lys Ala Ala Lys Met Gly Gly Gly Gly Arg Arg
                  100
                                      105
25
      (2) INFORMATION FOR SEQ ID NO:379:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1623 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1623
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025544
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:379:
      atgggcattt ctaagatgat tcttctgaaa ttcacatatg ttgtttcttt ctggattact
                                                                              60
      tgcgtgatct tcaaaatgtc aagaacgata aaacgagctt tgttgggctc attcatcttc
                                                                             120
      attettgett cegetagtgt tgtggtagee gecattgtag gagecatega aggteacace
                                                                             180
40
      actgacatcg gatttettea aggtagtgtg cttggagtgg tggetggagt tateactgeg
                                                                             240
      qtccaactct ttqqaccqqt qctacataqt qatcaacctt tqtctaaqqt qqctttactc
                                                                             300
      aggagagtag tgaatgggaa agccattatg ggattggtta gaccttttgt tctcaaagca
                                                                             360
      tatcaatggc aaataatagc attggataca agttacatgg agagttcaaa cttatacgat
                                                                             420
      ttcaatcatq aaaaaaqqq actatcqaaq agctctattc agaacatccc qatgttctac
                                                                             480
45
      aaccqttcaq aacatcaaac aaaatcqaqt tqctcqattt qcttacaqqa ttqqqaaqaa
                                                                             540
      qqqqaaqtaq qaaqaaaqct aqcaaqatqt qqccacacat ttcatatqaa ttqcataqat
                                                                             600
      gagtqqttqc ttagacaaga aacttgcccc atttgaagaa gacaaaagaa gaagcacqaq
                                                                             660
      aatcatctgc ttaaaatcca atcgcatcga ctcagtcttc ttcttcttct tcacctacct
                                                                             720
      ttctctctct ctctcgtggt ccattcgcta ttctatactt cttgtcggcg ccggagagag
                                                                             780
50
      tgagatgact tggtcggttt tcagatccat aaatactcca acactcgacc tctccaccgc
                                                                             840
      acttegetee actegtacee cattggtege tgetggtgte ggetgegeaa cattegetgg
                                                                             900
      tgtttctctc ttcagaatgt cttctagatc tcctcctttc gcttccctct ctgtctctgc
                                                                             960
      ttcttctgtg aagaaggaag ttgtgtctac tgagaaagca ccagctgctt tgggacctta
                                                                            1020
      ctctcaggcc attaaagcca ataatctggt ttttctttca ggtgttcttg gacttatacc
                                                                            1080
55
      tgagactgga aagtttgttt cggagagcgt cgaagatcag actgagcagg tactcaaaaa
                                                                            1140
      catgggggag atattgaaag ctagtggtgc tgattattcc tcggtggtga agacaacaat
                                                                            1200
      catgttggct gatttggctg acttcaagac agtgaacgag atatatgcca aatacttccc
                                                                            1260
      ageteettet ceageacqat egacgtatea agttgcaget ttgcctctaa acgecaaqat
                                                                            1320
      cqaqattqaa tqtattqcaa cactctaqaa cactcaaatc aatctctacq aaqqqtattt
                                                                            1380
60
      tggtccggaa gatgtcaaaa caaataagag aaacaatggt ttttggggct taacccatat
                                                                            1440
      ttaacctact tttcattttq ctqcactacq aaaaatqttq cqqattatca accattaqqt
                                                                            1500
      tggttacaaa tcaaaagcct ctaatcaaga ataagttatt tagtgtcttg tccaccctca
                                                                            1560
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512

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tttgctqttq acgtatatgt attgattgta tacattattt tatttgagtq aaaccattat 1620
     (2) INFORMATION FOR SEQ ID NO:380:
         (i) SEQUENCE CHARACTERISTICS:
 5
              (A) LENGTH: 249 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
10
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..249
              (D) OTHER INFORMATION: / Ceres Seq. ID 2025545
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:380:
15
     Met Ser Gly Cys Leu Asp Lys Leu Ala Pro Phe Glu Glu Asp Lys
                                      10
     Arg Arg Ser Thr Arg Ile Ile Cys Leu Lys Ser Asn Arg Ile Asp Ser
                                  25
     Val Phe Phe Phe Phe Thr Tyr Leu Ser Leu Ser Leu Ser Trp Ser
20
                              40
     Ile Arg Tyr Ser Ile Leu Leu Val Gly Ala Gly Glu Ser Glu Met Thr
                          55
     Trp Ser Val Phe Arg Ser Ile Asn Thr Pro Thr Leu Asp Leu Ser Thr
                       70
                               75
25
     Ala Leu Arg Ser Thr Arg Thr Pro Leu Val Ala Ala Gly Val Gly Cys
                 85
                                  90
     Ala Thr Phe Ala Gly Val Ser Leu Phe Arg Met Ser Ser Arg Ser Pro
               100 105 110
     Pro Phe Ala Ser Leu Ser Val Ser Ala Ser Ser Val Lys Lys Glu Val
30
                              120
                                                125
     Val Ser Thr Glu Lys Ala Pro Ala Ala Leu Gly Pro Tyr Ser Gln Ala
                          135
                                            140
     Ile Lys Ala Asn Asn Leu Val Phe Lev Ser Gly Val Leu Gly Leu Ile
                      150
                                         155
35
     Pro Glu Thr Gly Lys Phe Val Ser Glu Ser Val Glu Asp Gln Thr Glu
                                     170
     Gln Val Leu Lys Asn Met Gly Glu Ile Leu Lys Ala Ser Gly Ala Asp
                                  185 190
     Tyr Ser Ser Val Val Lys Thr Thr Ile Met Leu Ala Asp Leu Ala Asp
40
                              200
     Phe Lys Thr Val Asn Glu Ile Tyr Ala Lys Tyr Phe Pro Ala Pro Ser
                          215 220
     Pro Ala Arg Ser Thr Tyr Gln Val Ala Ala Leu Pro Leu Asn Ala Lys
                      230
                                         235
     Ile Glu Ile Glu Cys Ile Ala Thr Leu
45
                   245
     (2) INFORMATION FOR SEQ ID NO:381:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 211 amino acids
50
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
55
              (A) NAME/KEY: peptide
               (B) LOCATION: 1..211
              (D) OTHER INFORMATION: / Ceres Seq. ID 2025546
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:381:
     Met Gly Ile Ser Lys Met Ile Leu Leu Lys Phe Thr Tyr Val Val Ser
                5
60
                               10
     Phe Trp Ile Thr Cys Val Ile Phe Lys Met Ser Arg Thr Ile Lys Arg
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513 Ala Leu Leu Gly Ser Phe Ile Phe Ile Leu Ala Ser Ala Ser Val Val 40 Val Ala Ala Ile Val Gly Ala Ile Glu Gly His Thr Thr Asp Ile Gly 55 5 Phe Leu Gln Gly Ser Val Leu Gly Val Val Ala Gly Val Ile Thr Ala 75 Val Gln Leu Phe Gly Pro Val Leu His Ser Asp Gln Pro Leu Ser Lys 8.5 Val Ala Leu Leu Arg Arg Val Val Asn Gly Lys Ala Ile Met Gly Leu 10 105 Val Arg Pro Phe Val Leu Lys Ala Tyr Gln Trp Gln Ile Ile Ala Leu 120 Asp Thr Ser Tyr Met Glu Ser Ser Asn Leu Tyr Asp Phe Asn His Glu 135 140 15 Lys Lys Gly Leu Ser Lys Ser Ser Ile Gln Asn Ile Pro Met Phe Tyr 150 155 Asn Arg Ser Glu His Gln Thr Lys Ser Ser Cys Ser Ile Cys Leu Gln 170 Asp Trp Glu Glu Gly Glu Val Gly Arg Lys Leu Ala Arg Cys Gly His 20 185 190 Thr Phe His Met Asn Cys Ile Asp Glu Trp Leu Leu Arg Gln Glu Thr 195 200 Cys Pro Ile 210 25 (2) INFORMATION FOR SEQ ID NO:382: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 206 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 30 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..206 35 (D) OTHER INFORMATION: / Ceres Seq. ID 2025547 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:382: Met Ile Leu Leu Lys Phe Thr Tyr Val Val Ser Phe Trp Ile Thr Cys 10 Val Ile Phe Lys Met Ser Arg Thr Ile Lys Arg Ala Leu Leu Gly Ser 40 25 Phe Ile Phe Ile Leu Ala Ser Ala Ser Val Val Val Ala Ala Ile Val 40 Gly Ala Ile Glu Gly His Thr Thr Asp Ile Gly Phe Leu Gln Gly Ser 55 45 Val Leu Gly Val Val Ala Gly Val Ile Thr Ala Val Gln Leu Phe Gly 7.0 7.5 Pro Val Leu His Ser Asp Gln Pro Leu Ser Lys Val Ala Leu Leu Arg 90 8.5 Arg Val Val Asn Gly Lys Ala Ile Met Gly Leu Val Arg Pro Phe Val 50 105 Leu Lys Ala Tyr Gln Trp Gln Ile Ile Ala Leu Asp Thr Ser Tyr Met 120 Glu Ser Ser Asn Leu Tyr Asp Phe Asn His Glu Lys Lys Gly Leu Ser 135 140 55 Lys Ser Ser Ile Gln Asn Ile Pro Met Phe Tyr Asn Arg Ser Glu His 155 150 Gln Thr Lys Ser Ser Cys Ser Ile Cys Leu Gln Asp Trp Glu Glu Gly 165 170 Glu Val Gly Arg Lys Leu Ala Arg Cys Gly His Thr Phe His Met Asn

185

Cys Ile Asp Glu Trp Leu Leu Arg Gln Glu Thr Cys Pro Ile 195 200 205

180

514

```
(2) INFORMATION FOR SEQ ID NO:383:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 542 base pairs
                (B) TYPE: nucleic acid
 5
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
10
                (B) LOCATION: 1..542
                (D) OTHER INFORMATION: / Ceres Seq. ID 2026207
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:383:
      tcaccaaaat ctcttctctc tcttccattg tcttcctcta atggaaccaa ccgagaaatc
                                                                              60
      tatgttacta gaaactacta gtaccacgaa gatggagacc aaatacgaag atatgttacc
                                                                             120
15
      agttatggcg gagaagatgg atgttgaaga gtttgtatca gagttatgca aaggtttcag
                                                                             180
      tttgcttgcg gatccagaga gacatctcat cacagctgag tctctaagac gaaactcagg
                                                                             240
      gatacttgga attgaaggta tgagcaagga agatgctcaa ggaatggtta gagaaggaga
                                                                             300
      cctcgatgga gatggtgctc ttaaccaaac cgaattctgc gttctcatgg ttcggttaag
                                                                            360
      ccctgagatg atggaagacg ccgaaacttg gttggagaaa gcactcaccc aagaactatg
20
      taatcacaat ctctcttcta tgccttgaat gatcgtcctc ctttcttgtg tattctcttt
                                                                            480
      ttacttctac aaaacctata aatgtttctg aatataaaat gaagtaatga tttcttgatc
                                                                            540
      (2) INFORMATION FOR SEQ ID NO:384:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 148 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..148
                (D) OTHER INFORMATION: / Ceres Seq. ID 2026208
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:384:
35
      His Gln Asn Leu Phe Ser Leu Phe His Cys Leu Pro Leu Met Glu Pro
                                          10
      Thr Glu Lys Ser Met Leu Leu Glu Thr Thr Ser Thr Thr Lys Met Glu
                  20
                                      25
      Thr Lys Tyr Glu Asp Met Leu Pro Val Met Ala Glu Lys Met Asp Val
40
                                  4.0
      Glu Glu Phe Val Ser Glu Leu Cys Lys Gly Phe Ser Leu Leu Ala Asp
                              55
      Pro Glu Arg His Leu Ile Thr Ala Glu Ser Leu Arg Arg Asn Ser Gly
                          70
                                              7.5
45
      Ile Leu Gly Ile Glu Gly Met Ser Lys Glu Asp Ala Gln Gly Met Val
                                          90
      Arg Glu Gly Asp Leu Asp Gly Asp Gly Ala Leu Asn Gln Thr Glu Phe
                  100
                                      105
      Cys Val Leu Met Val Arg Leu Ser Pro Glu Met Met Glu Asp Ala Glu
50
                                  120
      Thr Trp Leu Glu Lys Ala Leu Thr Gln Glu Leu Cys Asn His Asn Leu
          130
                              135
                                                   140
      Ser Ser Met Pro
      145
55
      (2) INFORMATION FOR SEQ ID NO:385:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 135 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
60
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
```

(ix) FEATURE:

```
(A) NAME/KEY: peptide
               (B) LOCATION: 1..135
                (D) OTHER INFORMATION: / Ceres Seq. ID 2026209
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:385:
 5
      Met Glu Pro Thr Glu Lys Ser Met Leu Leu Glu Thr Thr Ser Thr Thr
                                         10
      Lys Met Glu Thr Lys Tyr Glu Asp Met Leu Pro Val Met Ala Glu Lys
                                     25
      Met Asp Val Glu Glu Phe Val Ser Glu Leu Cys Lys Gly Phe Ser Leu
10
                                 40
      Leu Ala Asp Pro Glu Arg His Leu Ile Thr Ala Glu Ser Leu Arg Arg
                             55
      Asn Ser Gly Ile Leu Gly Ile Glu Gly Met Ser Lys Glu Asp Ala Gln
                         7 Ó
15
      Gly Met Val Arg Glu Gly Asp Leu Asp Gly Asp Gly Ala Leu Asn Gln
                                         90
      Thr Glu Phe Cys Val Leu Met Val Arg Leu Ser Pro Glu Met Met Glu
                                    105
      Asp Ala Glu Thr Trp Leu Glu Lys Ala Leu Thr Gln Glu Leu Cys Asn
20
             115
                                120
      His Asn Leu Ser Ser Met Pro
         130
                     135
      (2) INFORMATION FOR SEQ ID NO:386:
          (i) SEQUENCE CHARACTERISTICS:
25
               (A) LENGTH: 128 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..128
               (D) OTHER INFORMATION: / Ceres Seq. ID 202 210
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:386:
35
     Met Leu Leu Glu Thr Thr Ser Thr Thr Lys Met Glu Thr Lys Tyr Glu
                                        10
      Asp Met Leu Pro Val Met Ala Glu Lys Met Asp Val Glu Glu Phe Val
                                    25
      Ser Glu Leu Cys Lys Gly Phe Ser Leu Leu Ala Asp Pro Glu Arg His
40
                                 40
      Leu Ile Thr Ala Glu Ser Leu Arg Arg Asn Ser Gly Ile Leu Gly Ile
                             55
      Glu Gly Met Ser Lys Glu Asp Ala Gln Gly Met Val Arg Glu Gly Asp
                         70
                                             75
45
     Leu Asp Gly Asp Gly Ala Leu Asn Gln Thr Glu Phe Cys Val Leu Met
                                         90
      Val Arg Leu Ser Pro Glu Met Met Glu Asp Ala Glu Thr Trp Leu Glu
                100 105 110
      Lys Ala Leu Thr Gln Glu Leu Cys Asn His Asn Leu Ser Ser Met Pro
50
                             120
             115
      (2) INFORMATION FOR SEQ ID NO:387:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 552 base pairs
               (B) TYPE: nucleic acid
55
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
               (A) NAME/KEY: -
60
               (B) LOCATION: 1..552
               (D) OTHER INFORMATION: / Ceres Seq. ID 2026982
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:387:
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WO 00/40695 PCT/US00/00466 516 atgaagtata aaaacgtttt tcttaaatta tggatgtctt ctctttatat agtgcccact 60 tecteattag cattetacag tetgeacaet tateaettet ttegaaatet etetetatee 120 180 ttcattgcca cagggatggg tccagtcacc gtggaggcac gcacgtgtga gtcaaaaaqc 240 cataggttca agggtccatg tgtgagcaca cacaactgtg caaacgtgtg ccacaacgaa 300 ggcttcggcg gaggtaaatg ccgtggattc cgtcgtcgtt gctactgcac aagacactqc 360 tgatccatcc attctcatga ctcaaatctt cgatccatcg tcagtgtgtt acttctttct 420 tatctaaatc ttccqtacqq taccatqtcq taccqtacat qaqtqttttc tcqaataaqt 480 cattggtttg tgtgtttccg gttttaatgt aatgttaaat caattaatgg cttttaatat 540 attgtattat gg (2) INFORMATION FOR SEQ ID NO:388: (i) SEOUENCE CHARACTERISTICS: (A) LENGTH: 120 amino acids

- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (ix) FEATURE:

5

10

15

20

40

45

- (A) NAME/KEY: peptide
- (B) LOCATION: 1..120
  - (D) OTHER INFORMATION: / Ceres Seq. ID 2026983
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:388:

Met Lys Tyr Lys Asn Val Phe Leu Lys Leu Trp Met Ser Ser Leu Tyr 1 5 10 15

25 Ile Val Pro Thr Ser Ser Leu Ala Phe Tyr Ser Leu His Thr Tyr His 20 25 30

Phe Phe Arg Asn Leu Ser Leu Ser Leu Ser Lys Met Lys Leu Ser Val
35 40 45

Arg Phe Ile Ser Ala Ala Leu Leu Phe Met Val Phe Ile Ala Thr 30 50 55 60 Gly Met Gly Pro Val Thr Val Glu Ala Arg Thr Cys Glu Ser Lys Ser

65 70 75 80
His Arg Phe Lys Gly Pro Cys Val Ser Thr His As Cys Ala Asn Val

85 90 95 35 Cys His Asn Glu Gly Phe Gly Gly Gly Lys Cys Arg Gly Phe Arg Arg 100 105 110

Arg Cys Tyr Cys Thr Arg His Cys 115 120

- (2) INFORMATION FOR SEQ ID NO:389:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 109 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: peptide
    - (B) LOCATION: 1..109
    - (D) OTHER INFORMATION: / Ceres Seq. ID 2026984

50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:389:

Met Ser Ser Leu Tyr Ile Val Pro Thr Ser Ser Leu Ala Phe Tyr Ser
1 5 10 15
Leu His Thr Tyr His Phe Phe Arg Asn Leu Ser Leu Ser Leu Ser Lys

55 Met Lys Leu Ser Val Arg Phe Ile Ser Ala Ala Leu Leu Leu Phe Met 35 40 45

Val Phe Ile Ala Thr Gly Met Gly Pro Val Thr Val Glu Ala Arg Thr 50 55 60

Cys Glu Ser Lys Ser His Arg Phe Lys Gly Pro Cys Val Ser Thr His 60 65 70 75 80

Asn Cys Ala Asn Val Cys His Asn Glu Gly Phe Gly Gly Gly Lys Cys 85 90 95

517 Arg Gly Phe Arg Arg Cys Tyr Cys Thr Arg His Cys 105 100 (2) INFORMATION FOR SEQ ID NO:390: (i) SEQUENCE CHARACTERISTICS: 5 (A) LENGTH: 77 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 10 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..77 (D) OTHER INFORMATION: / Ceres Seq. ID 2026985 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:390: 15 Met Lys Leu Ser Val Arg Phe Ile Ser Ala Ala Leu Leu Leu Phe Met 10 Val Phe Ile Ala Thr Gly Met Gly Pro Val Thr Val Glu Ala Arg Thr 25 20 Cys Glu Ser Lys Ser His Arg Phe Lys Gly Pro Cys Val Ser Thr His 20 40 Asn Cys Ala Asn Val Cys His Asn Glu Gly Phe Gly Gly Lys Cys 55 Arg Gly Phe Arg Arg Cys Tyr Cys Thr Arg His Cys 70 25 (2) INFORMATION FOR SEQ ID NO:391: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 66 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..66 35 (D) OTHER INFORMATION: / Ceres Seq. ID 2026986 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:391: Met Asp Val Phe Ser Leu Tyr Ser Ala His Phe Leu Ile Ser Ile Leu 10 Gln Ser Ala His Leu Ser Leu Leu Ser Lys Ser Leu Ser Ile Pro Leu 40 20 25 Lys Asn Glu Ala Leu Cys Ala Phe Tyr Leu Arg Cys Ser Ser Leu Val 40 His Gly Ile His Cys His Arg Asp Gly Ser Ser His Arg Gly Gly Thr 50 55 45 His Val (2) INFORMATION FOR SEQ ID NO:392: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1760 base pairs 50 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 55 (A) NAME/KEY: -(B) LOCATION: 1..1760 (D) OTHER INFORMATION: / Ceres Seq. ID 2027300 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:392: ctcgtttcat gagagaaaca aaaaatctta tctttctgac aacggtggtg gtgtgtgatg 60 60 ttgtcagcca tggcaaagac agctacaacg atctctctct cttttcttct tcttctgtgt 120 ctgtctttaa atagagtctc tttgaaacga agcagctttg tcctgttctg atatccatgg 180 ctttaccttc qtqcttaaag actggagctt taatgtctcc ggccactgga tttaattttt 240

518

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ccqqcaqtct tatqaaqtca qacaqcqqct tcqccqttcc gacaaaqtta caqaqtactc
                                                                             300
      qaaaaqqtqa tcqaqaqaqa ttqaqaqtcc aaqctatttt caqtttccct cctqcqtttc
                                                                             360
     tqacaaqaaa cggtcgagct gagaaacaga aacagctcaa acaagaactt cttgaagcca
                                                                             420
     ttgagcctct tgaacgtggt gctacggcct cgcctgatga ccagcttcgg attgatcagt
                                                                             480
5
     tagcgcgtaa agtggaagca gttaatccca ccaaggagcc tttgaagtct gatttggtca
                                                                             540
      atqqaaaatq qqaqctcatt tatacaacct ctqcttcgat tttqcaqqca aaqaaaccaa
                                                                             600
      ggttcttaag atcaataacc aactaccaat ctatcaatgt ggatacactt aaggtgcaaa
                                                                             660
      acatggagac ttggcctttc tataactcgg taactggaga cataaaaccc ctcaattcga
                                                                             720
      aqaaqqttqc tqtqaaactc caaqtqttta aaattctcqq atttattcct ataaaaqcac
                                                                             780
10
      ctgatagcgc ccgcggtgaa cttgagatta cctatgtgga cgaggaacta cggttatcaa
                                                                             840
      gagatettte atteatgtet catgtetggt tecatgacet ttetacattg tttetgttga
                                                                             900
      tttatqaatc ttcaataaca ctacaaqtta aaccaattag aagtgagttg qttatqqtcc
                                                                             960
      ggtacaaaga tcacaaccgg agaaggattg aagagtggtc aaagcgatgg tcagagttgg
                                                                            1020
      ttggttccgt cgaaacagga agaaagagtc tctcagaaac gacccagcca ctctttaaaa
                                                                            1080
     ctccattgga aaagctttgt cctgttttga tatccatggc tttaccttgg tgcttaaaga
15
                                                                            1140
      ctggagtttt aacgtctccg gccgccggat ttaatcatcc atcggacagt ggcttcgccg
                                                                            1200
      ttccgacgaa gttactgagt attcgaaaag gtgatcggga aagattgaga atccaagctg
                                                                            1260
      ttttcaqctt tcctccaaga aacggtggag ccgagaagcg gaaacagctc aaacatgaac
                                                                            1320
      ttgtcgaage cattgageet ettgaaegtg gtgeeactge etegeetgat gateagette
                                                                           1380
20
      tgattgatca gttagcacgt aaggtggaag cagtaaaccc aaccaaggag cctctgaagt
                                                                           1440
      ctgatttgat caatgggaaa tgggagctca tttacacaac atctgctgcg attttgcaag
                                                                           1500
      caaagaaacc aaggttotta agatogttaa otaactacca atgtatcaat atggatacac
                                                                           1560
      taaaggtgca aagaatggag acttggcctt tctataactc ggtaactgga gacttgacac
      ccctcaactc gaagacggtt gctgtgaaac ttcaagtgtt taaaattctc ggctttattc
                                                                           1680
25
      cggtaaaagc acctgatggt actgcacgcg gtgaactaga gattacctat gtggacgagg
                                                                            1740
      aactacqcqa caaactttqa
      (2) INFORMATION FOR SEQ ID NO:393:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 527 amino acids
30
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: pept_de
          (ix) FEATURE:
35
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..527
                (D) OTHER INFORMATION: / Ceres Seq. ID 2027301
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:393:
      Met Ala Leu Pro Ser Cys Leu Lys Thr Gly Ala Leu Met Ser Pro Ala
40
                                          10
      Thr Gly Phe Asn Phe Ser Gly Ser Leu Met Lys Ser Asp Ser Gly Phe
                                      25
      Ala Val Pro Thr Lys Leu Gln Ser Thr Arg Lys Gly Asp Arg Glu Arg
                                  40
45
      Leu Arg Val Gln Ala Ile Phe Ser Phe Pro Pro Ala Phe Leu Thr Arg
                              5.5
      Asn Gly Arg Ala Glu Lys Gln Lys Gln Leu Lys Gln Glu Leu Leu Glu
                                              75
      Ala Ile Glu Pro Leu Glu Arg Gly Ala Thr Ala Ser Pro Asp Asp Gln
50
                                          90
      Leu Arg Ile Asp Gln Leu Ala Arg Lys Val Glu Ala Val Asn Pro Thr
                                      105
      Lys Glu Pro Leu Lys Ser Asp Leu Val Asn Gly Lys Trp Glu Leu Ile
                                  120
55
      Tyr Thr Thr Ser Ala Ser Ile Leu Gln Ala Lys Lys Pro Arg Phe Leu
                              135
      Arg Ser Ile Thr Asn Tyr Gln Ser Ile Asn Val Asp Thr Leu Lys Val
                          150
                                              155
      Gln Asn Met Glu Thr Trp Pro Phe Tyr Asn Ser Val Thr Gly Asp Ile
60
                                          170
                      165
                                                              175
      Lys Pro Leu Asn Ser Lys Lys Val Ala Val Lys Leu Gln Val Phe Lys
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185

519 Ile Leu Gly Phe Ile Pro Ile Lys Ala Pro Asp Ser Ala Arg Gly Glu 200 Leu Glu Ile Thr Tyr Val Asp Glu Glu Leu Arg Leu Ser Arg Asp Leu 215 220 5 Ser Phe Met Ser His Val Trp Phe His Asp Leu Ser Thr Leu Phe Leu 230 235 Leu Ile Tyr Glu Ser Ser Ile Thr Leu Gln Val Lys Pro Ile Arg Ser 250 245 Glu Leu Val Met Val Arg Tyr Lys Asp His Asn Arg Arg Ile Glu 10 265 Glu Trp Ser Lys Arg Trp Ser Glu Leu Val Gly Ser Val Glu Thr Gly 280 Arg Lys Ser Leu Ser Glu Thr Thr Gln Pro Leu Phe Lys Thr Pro Leu 295 15 Glu Lys Leu Cys Pro Val Leu Ile Ser Met Ala Leu Pro Trp Cys Leu 310 315 Lys Thr Gly Val Leu Thr Ser Pro Ala Ala Gly Phe Asn His Pro Ser 325 330 Asp Ser Gly Phe Ala Val Pro Thr Lys Leu Leu Ser Ile Arg Lys Gly 20 340 345 350 Asp Arg Glu Arg Leu Arg Ile Gln Ala Val Phe Ser Phe Pro Pro Arg 360 Asn Gly Gly Ala Glu Lys Arg Lys Gln Leu Lys His Glu Leu Val Glu 375 25 Ala Ile Glu Pro Leu Glu Arg Gly Ala Thr Ala Ser Pro Asp Asp Gln 390 395 Leu Leu Ile Asp Gln Leu Ala Arg Lys Val Glu Ala Val Asn Pro Thr 410 Lys Glu Pro Leu Lys Ser Asp Leu Ile Asn Gly Lys Trp Glu Leu Ile 30 425 Tyr Thr Thr Ser Ala Ala Ile Leu Gln Ala Lys Lys Pro Arg Phe Leu 440 Arg Ser Leu Thr Asn Tyr Gln Cys Ile Asn Met Asp Thr Leu Lys Val 455 460 35 Gln Arg Met Glu Thr Trp Pro Phe Tyr Asn Ser Val Thr Gly Asp Leu 475 470 Thr Pro Leu Asn Ser Lys Thr Val Ala Val Lys Leu Gln Val Phe Lys 485 490 Ile Leu Gly Phe Ile Pro Val Lys Ala Pro Asp Gly Thr Ala Arg Gly 40 500 505 510 Glu Leu Glu Ile Thr Tyr Val Asp Glu Glu Leu Arg Asp Lys Leu 520 515 (2) INFORMATION FOR SEQ ID NO:394: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 515 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 50 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..515 (D) OTHER INFORMATION: / Ceres Seq. ID 2027302 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:394: 55 Met Ser Pro Ala Thr Gly Phe Asn Phe Ser Gly Ser Leu Met Lys Ser 10 Asp Ser Gly Phe Ala Val Pro Thr Lys Leu Gln Ser Thr Arg Lys Gly 25 Asp Arg Glu Arg Leu Arg Val Gln Ala Ile Phe Ser Phe Pro Pro Ala 40 60 45 Phe Leu Thr Arg Asn Gly Arg Ala Glu Lys Gln Lys Gln Leu Lys Gln 55

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		520														
	Glu 65	Leu	Leu	Glu	Ala	Ile 70	Glu	Pro	Leu	Glu	Arg 75	Gly	Ala	Thr	Ala	Ser 80
		Asp	Asp	Gln	Leu 85		Ile	Asp	Gln	Leu 90		Arg	Lys	Val	Glu 95	
5	Val	Asn	Pro	Thr 100		Glu	Pro	Leu	Lys 105		Asp	Leu	Val	Asn 110		Lys
	Trp	Glu	Leu 115		Tyr	Thr	Thr	Ser 120		Ser	Ile	Leu	Gln 125		Lys	Lys
10	Pro	Arg 130		Leu	Arg	Ser	Ile 135		Asn	Tyr	Gln	Ser 140	Ile	Asn	Val	Asp
	Thr 145	Leu	Lys	Val	Gln	Asn 150	Met	Glu	Thr	Trp	Pro 155	Phe	Tyr	Asn	Ser	Val 160
		Gly	Asp	Ile	Lys 165		Leu	Asn	Ser	Lys 170		Val	Ala	Val	Lys 175	
15	Gln	Val	Phe	Lys 180		Leu	Gly	Phe	Ile 185	Pro	Ile	Lys	Ala	Pro 190		Ser
	Ala	Arg	Gly 195	Glu	Leu	Glu	Ile	Thr 200	Tyr	Val	Asp	Glu	Glu 205	Leu	Arg	Leu
20	Ser	Arg 210	Asp	Leu	Ser	Phe	Met 215	Ser	His	Val	Trp	Phe 220	His	Asp	Leu	Ser
	Thr 225	Leu	Phe	Leu	Leu	Ile 230	Tyr	Glu	Ser	Ser	Ile 235	Thr	Leu	Gln	Val	Lys 240
	Pro	Ile	Arg	Ser	Glu 245	Leu	Val	Met	Val	Arg 250	Tyr	Lys	Asp	His	Asn 255	Arg
25	Arg	Arg	Ile	Glu 260	Glu	Trp	Ser	Lys	Arg 265	Trp	Ser	Glu	Leu	Val 270	Gly	Ser
	Val	Glu	Thr 275	Gly	Arg	Lys	Ser	Leu 280	Ser	Glu	Thr	Thr	Gln 285	Pro	Leu	Phe
30	Lys	Thr 290	Pro	Leu	Glu	Lys	Leu 295	Cys	Pro	Val	Leu	Ile 300	Ser	Met	Ala	Leu
	Pro 305	Trp	Cys	Leu	Lys	Thr 310	Gly	Val	Leu	Thr	Ser 315	Pro	Ala	Ala	Gly	Phe 320
	Asn	His	Pro	Ser	Asp 325	Ser	Gly	Phe	Ala	Val 330	Pro	Thr	Lys	Leu	Leu 335	Ser
35		_	-	340	=			_	345				Ala	350		
			355	_				360				_	Gln 365		_	
40		370					375					380	Ala			
	385	_	_			390		_			395	_	Lys			400
					405					410			Ile		415	_
45	_			420	_				425				Gln	430	_	_
			435					440					Ile 445			
50		450	-				455					460	Tyr			
	465					470					475		Ala			480
				_	485					490			Ala		495	_
55	Thr	Ala	Arg	Gly 500	Glu	Leu	Glu	Ile	Thr 505	Tyr	Val	Asp	Glu	Glu 510	Leu	Arg
	Asp	Lys	515													
60	(2)							NO:3 ISTI								
-		`-		-				mino		<i>d</i> ~						

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 502 amino acids (B) TYPE: amino acid

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(C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..502
               (D) OTHER INFORMATION: / Ceres Seq. ID 2027303
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:395:
     Met Lys Ser Asp Ser Gly Phe Ala Val Pro Thr Lys Leu Gln Ser Thr
10
                                        10
     Arg Lys Gly Asp Arg Glu Arg Leu Arg Val Gln Ala Ile Phe Ser Phe
                                    25
     Pro Pro Ala Phe Leu Thr Arg Asn Gly Arg Ala Glu Lys Gln Lys Gln
     Leu Lys Gln Glu Leu Leu Glu Ala Ile Glu Pro Leu Glu Arq Gly Ala
15
     Thr Ala Ser Pro Asp Asp Gln Leu Arg Ile Asp Gln Leu Ala Arg Lys
                         70
     Val Glu Ala Val Asn Pro Thr Lys Glu Pro Leu Lys Ser Asp Leu Val
20
                                        90
     Asn Gly Lys Trp Glu Leu Ile Tyr Thr Thr Ser Ala Ser Ile Leu Gln
                100 105
     Ala Lys Lys Pro Arg Phe Leu Arg Ser Ile Thr Asn Tyr Gln Ser Ile
                                120 125
25
     Asn Val Asp Thr Leu Lys Val Gln Asn Met Glu Thr Trp Pro Phe Tyr
                            135
                                                140
     Asn Ser Val Thr Gly Asp Ile Lys Pro Leu Asn Ser Lys Lys Val Ala
                        1.50
                                           155
     Val Lys Leu Gln Val Phe Lys Ile Leu Gly Phe Ile Pro Ile Lys Ala
30
                    165
                                        170
     Pro Asp Ser Ala Arg Gly Glu Leu Glu Ile Thr Tyr Val Asp Glu Glu
                                    185
     Leu Arg Leu Ser Arg Asp Leu Ser Phe Met Ser His Val Trp Phe His
                                200
     Asp Leu Ser Thr Leu Phe Leu Leu Ile Tyr Glu Ser Ser Ile Thr Leu
35
                            215
                                                220
     Gln Val Lys Pro Ile Arg Ser Glu Leu Val Met Val Arg Tyr Lys Asp
                        230
                                            235
      His Asn Arg Arg Arg Ile Glu Glu Trp Ser Lys Arg Trp Ser Glu Leu
40
                                        250
                     245
     Val Gly Ser Val Glu Thr Gly Arg Lys Ser Leu Ser Glu Thr Thr Gln
                                    265
                 260
      Pro Leu Phe Lys Thr Pro Leu Glu Lys Leu Cys Pro Val Leu Ile Ser
                                280
                                                    285
45
     Met Ala Leu Pro Trp Cys Leu Lys Thr Gly Val Leu Thr Ser Pro Ala
                            295
                                                300
      Ala Gly Phe Asn His Pro Ser Asp Ser Gly Phe Ala Val Pro Thr Lys
                        310
                                            315
      Leu Leu Ser Ile Arg Lys Gly Asp Arg Glu Arg Leu Arg Ile Gln Ala
50
                     325
                                        330
      Val Phe Ser Phe Pro Pro Arg Asn Gly Gly Ala Glu Lys Arg Lys Gln
                 340
                                    345
      Leu Lys His Glu Leu Val Glu Ala Ile Glu Pro Leu Glu Arg Gly Ala
                                360
55
      Thr Ala Ser Pro Asp Asp Gln Leu Leu Ile Asp Gln Leu Ala Arg Lys
                            375
      Val Glu Ala Val Asn Pro Thr Lys Glu Pro Leu Lys Ser Asp Leu Ile
                         390
                                            395
      Asn Gly Lys Trp Glu Leu Ile Tyr Thr Thr Ser Ala Ala Ile Leu Gln
                         410 415
60
                     405
      Ala Lys Lys Pro Arg Phe Leu Arg Ser Leu Thr Asn Tyr Gln Cys Ile
                                    425
                 420
```

522 Asn Met Asp Thr Leu Lys Val Gln Arg Met Glu Thr Trp Pro Phe Tyr 440 Asn Ser Val Thr Gly Asp Leu Thr Pro Leu Asn Ser Lys Thr Val Ala 455 5 Val Lys Leu Gln Val Phe Lys Ile Leu Gly Phe Ile Pro Val Lys Ala 475 470 Pro Asp Gly Thr Ala Arg Gly Glu Leu Glu Ile Thr Tyr Val Asp Glu 490 485 Glu Leu Arg Asp Lys Leu 10 500 (2) INFORMATION FOR SEQ ID NO:396: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 539 base pairs (B) TYPE: nucleic acid 15 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -20 (B) LOCATION: 1..539 (D) OTHER INFORMATION: / Ceres Seq. ID 2027375 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:396: actgtgttcg ataataatgt cgacaggaga agcgatacca agagtcgccg tcqtcqtttt 60 catteteaac ggaaacteaa tettattagg tegeegeegt teeteaateg geaacteeac 120 25 tttcgctctt cccggtggcc acctcgaatt cggagagagc tttgaagaat gtgcaqcqaq 180 agaagtaatq qaqqaaacaq qtctaaaqat tgaaaagatg aagcttttga ctqttacaaa 240 caatqtcttc aaaqaaqcac caacqccatc acactacqtc tctqtttcga tacqtqcqqt gttggtggat ccaagtcaag aaccgaagaa tatggaacca gagaagtgtg aaggatggga ttqqtatqat tqqqaqaatc taccaaagcc tttgttttgg ccacttgaga aattgtttqq 30 aagtggtttc aatcetttca ctcatggtgg tggagactaa tagatgtaag agttaatgat tgatttggga ttgaatgttg cacaaattgg gcatttggtc tagtggtatg atctygctt (2) INFORMATION FOR SEO ID NO:397: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 152 amino acids 35 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 40 (A) NAME/KEY: peptide (B) LOCATION: 1..152 (D) OTHER INFORMATION: / Ceres Seq. ID 2027376 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:397: Leu Cys Ser Ile Ile Met Ser Thr Gly Glu Ala Ile Pro Arg Val Ala 45 10 Val Val Phe Ile Leu Asn Gly Asn Ser Ile Leu Leu Gly Arg Arg 20 25 Arg Ser Ser Ile Gly Asn Ser Thr Phe Ala Leu Pro Gly Gly His Leu 40 50 Glu Phe Gly Glu Ser Phe Glu Glu Cys Ala Ala Arg Glu Val Met Glu 55 Glu Thr Gly Leu Lys Ile Glu Lys Met Lys Leu Leu Thr Val Thr Asn 70 Asn Val Phe Lys Glu Ala Pro Thr Pro Ser His Tyr Val Ser Val Ser 55 90 Ile Arg Ala Val Leu Val Asp Pro Ser Gln Glu Pro Lys Asn Met Glu 105 Pro Glu Lys Cys Glu Gly Trp Asp Trp Tyr Asp Trp Glu Asn Leu Pro 120 60 Lys Pro Leu Phe Trp Pro Leu Glu Lys Leu Phe Gly Ser Gly Phe Asn 135

Pro Phe Thr His Gly Gly Asp

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145
                         150
      (2) INFORMATION FOR SEQ ID NO:398:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 147 amino acids
 5
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
10
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..147
               (D) OTHER INFORMATION: / Ceres Seq. ID 2027377
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:398:
     Met Ser Thr Gly Glu Ala Ile Pro Arg Val Ala Val Val Phe Ile
15
              5
                                         10
     Leu Asn Gly Asn Ser Ile Leu Leu Gly Arg Arg Arg Ser Ser Ile Gly
                                     25
     Asn Ser Thr Phe Ala Leu Pro Gly Gly His Leu Glu Phe Gly Glu Ser
                                 40
20
     Phe Glu Glu Cys Ala Ala Arg Glu Val Met Glu Glu Thr Gly Leu Lys
                             55
      Ile Glu Lys Met Lys Leu Leu Thr Val Thr Asn Asn Val Phe Lys Glu
                         70
                                            7.5
     Ala Pro Thr Pro Ser His Tyr Val Ser Val Ser Ile Arg Ala Val Leu
25
                                         90
                     85
     Val Asp Pro Ser Gln Glu Pro Lys Asn Met Glu Pro Glu Lys Cys Glu
                                    105 110
     Gly Trp Asp Trp Tyr Asp Trp Glu Asn Leu Pro Lys Pro Leu Phe Trp
             115 120 125
30
      Pro Leu Glu Lys Leu Phe Gly Ser Gly Phe Asn Pro Phe Thr His Gly
                            135
     Gly Gly Asp
      (2) INFORMATION FOR SEQ ID NO:399:
35
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 90 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..90
               (D) OTHER INFORMATION: / Ceres Seq. ID 2027378
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:399:
      Met Glu Glu Thr Gly Leu Lys Ile Glu Lys Met Lys Leu Leu Thr Val
                                        10
      Thr Asn Asn Val Phe Lys Glu Ala Pro Thr Pro Ser His Tyr Val Ser
                 20
                                     25
      Val Ser Ile Arg Ala Val Leu Val Asp Pro Ser Gln Glu Pro Lys Asn
50
                                 40
      Met Glu Pro Glu Lys Cys Glu Gly Trp Asp Trp Tyr Asp Trp Glu Asn
                             55
                                                60
      Leu Pro Lys Pro Leu Phe Trp Pro Leu Glu Lys Leu Phe Gly Ser Gly
55
                                            75
                         70
      Phe Asn Pro Phe Thr His Gly Gly Asp
                     85
      (2) INFORMATION FOR SEQ ID NO:400:
          (i) SEQUENCE CHARACTERISTICS:
60
                (A) LENGTH: 3983 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
```

524

(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: DNA (genomic)

(ix) FEATURE:

5

(A) NAME/KEY: -

(B) LOCATION: 1..3983

(D) OTHER INFORMATION: / Ceres Seq. ID 2028729

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:400:
     atgaacacga cgccgtttca ctcggatcct ccgccgtcga ggatccagcg taagctcgtt
                                                                            60
     gtcgaagttg ttgaagctcg taatattctc cctaaagatg gtcaaggaag ctctagcgct
                                                                            120
10
     tacgtcgttg tcgatttcga tgctcagaag aaacgaacct ccactaagtt ccgtgaccta
                                                                            180
     aaccctattt ggaacgagat gettgattte geegteteeg atcccaaaaa catggattae
                                                                            240
     gacgageteg atategaggt ttataacgat aaaagatttg gtaacggagg tggccggaag
                                                                            300
     aatcattttc tcqqtaqqqt taaqatctat qgaagccagt tctcqcgaag aqqtqaaqaa
                                                                            360
     ggtcttgtgt atttcccttt ggagaagaag agtgtgttca gctggattcg cggcgagatt
                                                                            420
15
     qqactcaaaa tctactatta cqacqaaqcc qccqacqaaq acacqqcqqq tqqaqqtqqa
                                                                            480
     ggacagcaac aacaacagca acagcaacaa tttcatccgc cgcaacaaga agccgatgaa
                                                                            540
     caacaacacc agcaacaatt tcatcctccg ccgcagcaga tgatgaatat accaccggag
                                                                            600
     aaacctaatq taqttqtqqt tqaaqaaqqt aqqqttttcq aatcqqctca qaqtcaqcqc
                                                                            660
     tatacaqaqa cacatcaqca acctccggtg gttattgttg aagaatcacc accgcaqcat
                                                                            720
20
     qtaatqcaaq qtccaaatqa taaccatcct caccgaaatq ataaccatcc tcaacqqcca
                                                                            780
     ccgtctccgc cgccacctcc atcggctggg gaagtacatt attatccacc ggaagtgagg
                                                                            840
     aagatgcaag taggaagacc tcccggcgga gatagaatta gggttacgaa gagaccaccq
                                                                            900
     aatggagatt attcacctag ggttatcaat agcaaaactg gaggaggaga gacgacgatg
                                                                            960
     gagaagaaga ctcatcatcc ttacaatctt gttgagccaa tgcagtatct cttcqttcqq
                                                                           1020
25
      attgtgaagg cgcgtggctt accacctaac gagagcgcgt atgttaaggt acggacgtcg
                                                                           1080
      aaccatttcg tcaggtctaa accggccgtt aaccggcccg gcgaatcggt tgattcaccq
                                                                           1140
      gagtggaatc aggtttttgc tcttggtcat aaccggtctg attccgctgt aactggtgcg
                                                                           1200
      actettgaga tetetgettg ggatgetteg teggagagtt tteteggagg agtttgtttt
                                                                           1260
      gatetetetg aggiteeggt tegigaeeeg eeggatagie egeitgetee teagiggiat
                                                                           1320
30
      eggetegaag geteeggege ggateagaae tetgggagaa titeeggtga eatteagete
                                                                           1380
      tctqtttqqa ttqqtactca qqtagatqaq qcatttccgg aggcttggag ctctqatqct
                                                                           1440
      ccgcatgtag ctcacacgcg ttctaaggtg tatcaatcgc cgaaactttg gtacttgaga
                                                                           1500
      gtgacggttc ttgaggcaca ggatttacac atagctccta atctcccgcc gttgactgcg
                                                                           1560
      cctgagattc gtgtgaaagc tcaattaggg tttcagtcgg cgcgtacaag aagaggctca
                                                                           1620
35
      atgaataacc acagtggttc gtttcattgg catgaggata tgatctttgt tgctqqaqaq
                                                                           1680
      ccgttggaag attgcttggt tctgatggtg gaagaccgga cgactaaaga agcaacactt
                                                                           1740
      ctaggacatg ccatgatece agtgagetee ategageage gaattgatga gegttttgtg
                                                                           1800
      ccgtcgaaat ggcacactct ggaaggagaa ggtggaggtg gaggtggagg aggaggacct
                                                                           1860
      ggaggtggtg gtggtggtgg accttattgt ggaaggatta gccttagact ttgtctcgaa
                                                                           1920
40
      ggtgggtatc atgtgcttga agaggcggcg catgtatgca gcgatttccg tccgacggct
                                                                           1980
      aagcagctat ggaaaccgcc gattggaata cttgagttgg ggattcttgg agctcgtggg
                                                                           2040
      ttgttgccga tgaaggcgaa aaacggaggg aaaggttcca ctgatgctta ttgtgttgct
                                                                           2100
      aagtacggga agaaatgggt caggactcga accataacag acagttttga cccgaggtgg
                                                                           2160
      cacgagcagt atacgtggca ggtttatgat ccttgcaccg tgctaactgt tggagtcttc
                                                                           2220
45
      gacaattgga ggatgttctc tgacgcctcc gatgatagac ctgacacacg gattgggaag
                                                                           2280
      atacggatec gggtgtcgac gttagagage aacaaagtgt acaccaatte atateetetg
                                                                           2340
      ttggttttgt tacctagcgg tatgaaaaaa atgggtgaaa ttgaagtggc agtccggttt
                                                                           2400
      geatgecegt etetgetgee tgatgtttgt geagettatg gacageeget tetgeetegg
                                                                           2460
      atgcactaca taaggcctct aggtgtagca caacaagatg cattaagagg ggccgccacg
                                                                           2520
50
      aaaatggtag cagcttggct ggctcgagca gaaccaccat tgggaccaga ggtagttcga
                                                                           2580
      2640
      agaattgttg gtgttttagc ttgggcagtg ggtttagcta agtggttgga taatatcagg
                                                                           2700
      cggtggagga atccagtgac gacggtgcta gtccatattc tatatctggt tcttgtttgg
                                                                           2760
      taccetgatt tggtagtece gactgeatte ttgtaegtgg tgatgategg agtttggtae
                                                                           2820
55
      taccggttta gacccaagat accggctggt atggatatcc gcttatcaca agctgaaacc
                                                                           2880
      qtcgatcctq atqaqctaqa tqaaqaattc qacaccatac caagctcaag qcgaccaqaa
                                                                           2940
      gtaatccgag ctaggtacga ccgattaagg atcttagcag tgagggttca gaccattcta
                                                                           3000
      ggagattttg cagcgcaagg agaacggatt caagcgttgg ttagctggag agatccgaga
                                                                           3060
      gcgacaaagc tgttcatagc aatctgtttg gtaatcacaa tagttctgta tgcagttcct
                                                                           3120
60
      qcqaaaatqq tqqcqqtggc tctaggagtt tcagatagtg tgccaacggc gaaacaagac
                                                                           3180
      acqaaaqaat ctctaaaqaa atcattttca tcgcttcgct tcgatttttc ttcaatqqct
                                                                           3240
      gtcgtcggcg ctccaatatc gtctccggcg gctcagctgc agacacaatt tctctccaat
                                                                           3300
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525

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cccattctcc cccqctttcq ccqqtctttc tccaccqgaa aatcaccagc aactttctcc
                                                                           3360
     gtcgtagcta tggctcccca gaaaaaggtg aacaaatatg atgccaagtg gaagaaacaa
                                                                           3420
     tggtacggag ctggattgtt tttcgaaggg agtgagcaaa taaacgttga tgttttcaag
                                                                           3480
     aagctggaga agcgaaaagt gttgagcaac gttgagaaat ctggcctgct gtcaaaagca
 5
     gaggggttgg gactcacatt gtcatctctt gagaagctta aagtcttctc caaagcagag
     qaccttqqtc ttctcaqtct ccttqaqaac ttaqctqqaa catcqcctqc qqtcttaqcc
     toggotgcat taccagotet cacqqotqct attgtagcog tggtgttgat cocqqatqac
     tcaactactc tagtggttgc tcaggcggtt ttggccggtg ctcttgcgct tacaggggtt
                                                                           3780
     gttttgttgg ttggttctgt tgttttggat ggacttcaag aagctgactg attctttctc
                                                                           3840
10
     tgtaaaccga acataaaccc atgtcttgtc caattgattt ttgtcagttg ctgatttata
                                                                           3900
     gctgtatggt tcagttgttt atggttagtc caagacataa gctgagtgat agaaagaagc
                                                                           3960
     tttataataa ttaaacaaat att
      (2) INFORMATION FOR SEQ ID NO:401:
           (i) SEQUENCE CHARACTERISTICS:
15
                (A) LENGTH: 1276 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
20
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..1276
                (D) OTHER INFORMATION: / Ceres Seq. ID 2028730
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:401:
25
     Met Asn Thr Thr Pro Phe His Ser Asp Pro Pro Pro Ser Arg Ile Gln
                                         10
                      5
     Arg Lys Leu Val Val Glu Val Glu Ala Arg Asn Ile Leu Pro Lys
                  2.0
                                      2.5
      Asp Gly Gln Gly Ser Ser Ser Ala Tyr Val Val Val Asp Phe Asp Ala
30
                                 40
      Gln Lys Lys Arg Thr Ser Thr Lys Phe Arg Asp Leu Asn Pro Ile Trp
                              55
      Asn Glu Met Leu Asp Phe Ala Val Ser Asp Pro Lys Asn Met Asp Tyr
                          70
                                              75
35
      Asp Glu Leu Asp Ile Glu Val Tyr Asn Asp Lys Arg Phe Gly Asn Gly
                      8.5
                                          90
      Gly Gly Arg Lys Asn His Phe Leu Gly Arg Val Lys Ile Tyr Gly Ser
                  100
                                      105
      Gln Phe Ser Arg Arg Gly Glu Glu Gly Leu Val Tyr Phe Pro Leu Glu
40
                                  120
      Lys Lys Ser Val Phe Ser Trp Ile Arg Gly Glu Ile Gly Leu Lys Ile
                              135
      Tyr Tyr Tyr Asp Glu Ala Ala Asp Glu Asp Thr Ala Gly Gly Gly
                                              155
                          150
45
      Gly Gln Gln Gln Gln Gln Gln Gln Phe His Pro Pro Gln Gln
                                          170
      Glu Ala Asp Glu Gln Gln His Gln Gln Gln Phe His Pro Pro Gln
                                      185
      Gln Met Met Asn Ile Pro Pro Glu Lys Pro Asn Val Val Val Glu
50
                                  200
      Glu Gly Arg Val Phe Glu Ser Ala Gln Ser Gln Arg Tyr Thr Glu Thr
                              215
                                                  220
      His Gln Gln Pro Pro Val Val Ile Val Glu Glu Ser Pro Pro Gln His
                          230
                                              235
55
      Val Met Gln Gly Pro Asn Asp Asn His Pro His Arg Asn Asp Asn His
                                          250
                      245
      Pro Gln Arg Pro Pro Ser Pro Pro Pro Pro Ser Ala Gly Glu Val
                                      265
                  260
      His Tyr Tyr Pro Pro Glu Val Arg Lys Met Gln Val Gly Arg Pro Pro
60
                                  280
      Gly Gly Asp Arg Ile Arg Val Thr Lys Arg Pro Pro Asn Gly Asp Tyr
```

295

290

526 Ser Pro Arg Val Ile Asn Ser Lys Thr Gly Gly Gly Glu Thr Thr Met Glu Lys Lys Thr His His Pro Tyr Asn Leu Val Glu Pro Met Gln Tyr 325 330 5 Leu Phe Val Arg Ile Val Lys Ala Arg Gly Leu Pro Pro Asn Glu Ser 345 Ala Tyr Val Lys Val Arg Thr Ser Asn His Phe Val Arg Ser Lys Pro 360 Ala Val Asn Arg Pro Gly Glu Ser Val Asp Ser Pro Glu Trp Asn Gln 10 380 375 Val Phe Ala Leu Gly His Asn Arg Ser Asp Ser Ala Val Thr Gly Ala 390 395 Thr Leu Glu Ile Ser Ala Trp Asp Ala Ser Ser Glu Ser Phe Leu Gly 410 15 Gly Val Cys Phe Asp Leu Ser Glu Val Pro Val Arg Asp Pro Pro Asp 425 420 Ser Pro Leu Ala Pro Gln Trp Tyr Arg Leu Glu Gly Ser Gly Ala Asp 440 Gln Asn Ser Gly Arg Ile Ser Gly Asp Ile Gln Leu Ser Val Trp Ile 20 455 460 Gly Thr Gln Val Asp Glu Ala Phe Pro Glu Ala Trp Ser Ser Asp Ala 465 470 475 Pro His Val Ala His Thr Arg Ser Lys Val Tyr Gln Ser Pro Lys Leu 485 490 495 25 Trp Tyr Leu Arg Val Thr Val Leu Glu Ala Gln Asp Leu His Ile Ala 500 505 510 Pro Asn Leu Pro Pro Leu Thr Ala Pro Glu Ile Arg Val Lys Ala Gln 520 Leu Gly Phe Gln Ser Ala Arg Thr Arg Arg Gly Ser Met Asn Asn His 30 540 530 535 Ser Gly Ser Phe His Trp His Glu Asp Met Ile Phe Val Ala Gly Glu 550 555 Pro Leu Glu Asp Cys Leu Val Leu Met Val Glu Asp Arg Thr Thr Lys 565 570 35 Glu Ala Thr Leu Leu Gly His Ala Met Ile Pro Val Ser Ser Ile Glu 585 Gln Arg Ile Asp Glu Arg Phe Val Pro Ser Lys Trp His Thr Leu Glu 600 Gly Glu Gly Gly Gly Gly Gly Gly Gly Pro Gly Gly Gly Gly 40 615 Gly Gly Gly Pro Tyr Cys Gly Arg Ile Ser Leu Arg Leu Cys Leu Glu 630 635 Gly Gly Tyr His Val Leu Glu Glu Ala Ala His Val Cys Ser Asp Phe 645 650 45 Arg Pro Thr Ala Lys Gln Leu Trp Lys Pro Pro Ile Gly Ile Leu Glu 665 Leu Gly Ile Leu Gly Ala Arg Gly Leu Leu Pro Met Lys Ala Lys Asn 680 Gly Gly Lys Gly Ser Thr Asp Ala Tyr Cys Val Ala Lys Tyr Gly Lys 50 695 700 Lys Trp Val Arg Thr Arg Thr Ile Thr Asp Ser Phe Asp Pro Arg Trp 715 710 His Glu Gln Tyr Thr Trp Gln Val Tyr Asp Pro Cys Thr Val Leu Thr 725 730 Val Gly Val Phe Asp Asn Trp Arg Met Phe Ser Asp Ala Ser Asp Asp 55 740 745 Arg Pro Asp Thr Arg Ile Gly Lys Ile Arg Ile Arg Val Ser Thr Leu 760 Glu Ser Asn Lys Val Tyr Thr Asn Ser Tyr Pro Leu Leu Val Leu Leu 60 775 Pro Ser Gly Met Lys Lys Met Gly Glu Ile Glu Val Ala Val Arg Phe

790

527 Ala Cys Pro Ser Leu Leu Pro Asp Val Cys Ala Ala Tyr Gly Gln Pro 805 810 Leu Leu Pro Arg Met His Tyr Ile Arg Pro Leu Gly Val Ala Gln Gln 825 5 Asp Ala Leu Arg Gly Ala Ala Thr Lys Met Val Ala Ala Trp Leu Ala 840 Arg Ala Glu Pro Pro Leu Gly Pro Glu Val Val Arg Tyr Met Leu Asp 855 Ala Asp Ser His Ala Trp Ser Met Arg Lys Ser Lys Ala Asn Trp Tyr 10 870 875 Arg Ile Val Gly Val Leu Ala Trp Ala Val Gly Leu Ala Lys Trp Leu 885 890 Asp Asn Ile Arg Arg Trp Arg Asn Pro Val Thr Thr Val Leu Val His 900 905 15 Ile Leu Tyr Leu Val Leu Val Trp Tyr Pro Asp Leu Val Val Pro Thr 920 Ala Phe Leu Tyr Val Val Met Ile Gly Val Trp Tyr Tyr Arg Phe Arg 935 940 Pro Lys Ile Pro Ala Gly Met Asp Ile Arg Leu Ser Gln Ala Glu Thr 20 950 955 Val Asp Pro Asp Glu Leu Asp Glu Glu Phe Asp Thr Ile Pro Ser Ser 970 975 965 Arg Arg Pro Glu Val Ile Arg Ala Arg Tyr Asp Arg Leu Arg Ile Leu 980 985 990 25 Ala Val Arg Val Gln Thr Ile Leu Gly Asp Phe Ala Ala Gln Gly Glu 995 1000 1005 Arg Ile Gln Ala Leu Val Ser Trp Arg Asp Pro Arg Ala Thr Lys Leu 1010 1015 1020 Phe Ile Ala Ile Cys Leu Val Ile Thr Ile Val Leu Tyr Ala Val Pro 30 1025 1030 1035 Ala Lys Met Val Ala Val Ala Leu Gly Val Ser Asp Ser Val Pro Thr 1045 1050 Ala Lys Gln Asp Thr Lys Glu Ser Leu Lys Lys Ser Phe Ser Ser Leu 1060 1065 1070 35 Arg Phe Asp Phe Ser Ser Met Ala Val Val Gly Ala Pro Ile Ser Ser 1075 1080 Pro Ala Ala Gln Leu Gln Thr Gln Phe Leu Ser Asn Pro Ile Leu Pro 1090 1095 1100 Arg Phe Arg Arg Ser Phe Ser Thr Gly Lys Ser Pro Ala Thr Phe Ser 40 1105 1110 1115 1120 Val Val Ala Met Ala Pro Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys 1125 1130 Trp Lys Lys Gln Trp Tyr Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu 1140 1145 1150 45 Gln Ile Asn Val Asp Val Phe Lys Lys Leu Glu Lys Arg Lys Val Leu 1155 1160 1165 Ser Asn Val Glu Lys Ser Gly Leu Leu Ser Lys Ala Glu Gly Leu Gly 1175 1180 Leu Thr Leu Ser Ser Leu Glu Lys Leu Lys Val Phe Ser Lys Ala Glu 50 1185 1190 1195 Asp Leu Gly Leu Leu Ser Leu Leu Glu Asn Leu Ala Gly Thr Ser Pro 1205 1210 Ala Val Leu Ala Ser Ala Ala Leu Pro Ala Leu Thr Ala Ala Ile Val 1225 1230 1220 55 Ala Val Val Leu Ile Pro Asp Asp Ser Thr Thr Leu Val Val Ala Gln 1240 1245 1235 Ala Val Leu Ala Gly Ala Leu Ala Leu Thr Gly Val Val Leu Leu Val 1255 1260 Gly Ser Val Val Leu Asp Gly Leu Gln Glu Ala Asp 60 1270 (2) INFORMATION FOR SEQ ID NO:402: (i) SEQUENCE CHARACTERISTICS:

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(A) LENGTH: 1210 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..1210
              (D) OTHER INFORMATION: / Ceres Seq. ID 2028731
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:402:
     Met Leu Asp Phe Ala Val Ser Asp Pro Lys Asn Met Asp Tyr Asp Glu
                                     10
     Leu Asp Ile Glu Val Tyr Asn Asp Lys Arg Phe Gly Asn Gly Gly
            20
                                  25
15
     Arg Lys Asn His Phe Leu Gly Arg Val Lys Ile Tyr Gly Ser Gln Phe
                               40
     Ser Arg Arg Gly Glu Glu Gly Leu Val Tyr Phe Pro Leu Glu Lys Lys
                           55
     Ser Val Phe Ser Trp Ile Arg Gly Glu Ile Gly Leu Lys Ile Tyr Tyr
20
                       70
     Tyr Asp Glu Ala Ala Asp Glu Asp Thr Ala Gly Gly Gly Gly Gln
                                     90
                   85
     Gln Gln Gln Gln Gln Gln Fhe His Pro Pro Gln Gln Glu Ala
                                  105
                100
25
     Asp Glu Gln Gln His Gln Gln Phe His Pro Pro Gln Gln Met
                              120
     Met Asn Ile Pro Pro Glu Lys Pro Asn Val Val Val Glu Glu Gly
                          135
                                             140
     Arg Val Phe Glu Ser Ala Gln Ser Gln Arg Tyr Thr Glu Thr His Gln
30
     145 150 155
     Gln Pro Pro Val Val Ile Val Glu Glu Ser Pro Pro Gln His Val Met
                    165 170 175
     Gln Gly Pro Asn Asp Asn His Pro His Arg Asn Asp Asn His Pro Gln
                     185
35
     Arg Pro Pro Ser Pro Pro Pro Pro Pro Ser Ala Gly Glu Val His Tyr
                              200
      Tyr Pro Pro Glu Val Arg Lys Met Gln Val Gly Arg Pro Pro Gly Gly
                           215
                                            220
      Asp Arg Ile Arg Val Thr Lys Arg Pro Pro Asn Gly Asp Tyr Ser Pro
40
                       230
                                          235
     Arg Val Ile Asn Ser Lys Thr Gly Gly Gly Glu Thr Thr Met Glu Lys
                                    250
      Lys Thr His His Pro Tyr Asn Leu Val Glu Pro Met Gln Tyr Leu Phe
                                   265
45
      Val Arg Ile Val Lys Ala Arg Gly Leu Pro Pro Asn Glu Ser Ala Tyr
                               280
      Val Lys Val Arg Thr Ser Asn His Phe Val Arg Ser Lys Pro Ala Val
                           295
      Asn Arg Pro Gly Glu Ser Val Asp Ser Pro Glu Trp Asn Gln Val Phe
50
                        310
                                          315
      Ala Leu Gly His Asn Arg Ser Asp Ser Ala Val Thr Gly Ala Thr Leu
                    325
                                       330
      Glu Ile Ser Ala Trp Asp Ala Ser Ser Glu Ser Phe Leu Gly Gly Val
                                   345
                340
      Cys Phe Asp Leu Ser Glu Val Pro Val Arg Asp Pro Pro Asp Ser Pro
55
                               360
      Leu Ala Pro Gln Trp Tyr Arg Leu Glu Gly Ser Gly Ala Asp Gln Asn
                           375
                                              380
      Ser Gly Arg Ile Ser Gly Asp Ile Gln Leu Ser Val Trp Ile Gly Thr
60
                        390
                                          395
      Gln Val Asp Glu Ala Phe Pro Glu Ala Trp Ser Ser Asp Ala Pro His
```

410

			His	420	_		_		425				_	430	_	-
5		-	Val 435					440		_			445			
	Leu	Pro 450	Pro	Leu	Thr	Ala	Pro 455	Glu	Ile	Arg	Val	Lys 460	Ala	Gln	Leu	Gly
	Phe 465	Gln	Ser	Ala	Arg	Thr 470	Arg	Arg	Gly	Ser	Met 475	Asn	Asn	His	Ser	Gly 480
10			His		485					490			_		495	
		_	Cys	500					505					510		
1 -			Leu 515	_				520					525			_
15		530	Glu	_			535		_	_		540			_	
	545		Gly			550					555			_	_	560
20	-		Tyr	-	565	-				570		-			575	-
	_		Val Lys	580					585		_			590	_	
25			595 Gly			_	-	600			-		605			-
25		610	Ser				615					620	_		_	_
	625	_	Thr			630					635	_	_	_	_	640
30		_	Thr		645					650					655	
		-	Asp	660			_	_	665	_				670		_
35			675 Arg		_	_		680					685	_	_	
	Asn	690 Lys	Val	Tyr	Thr	Asn	695 Ser	Tyr	Pro	Leu	Leu	700 Val	Leu	Leu	Pro	Ser
	705 Gly	Met	Lys	Lys	Met	710 Gly	Glu	Ile	Glu	Val	715 Ala	Val	Arg	Phe	Ala	720 Cys
40	Pro	Ser	Leu	Leu	725 Pro	Asp	Val	Cys	Ala	730 Ala	Tyr	Gly	Gln	Pro	735 Leu	Leu
	Pro	Arg	Met					Pro	_	Gly			Gln			Ala
45	Leu	_	755 Gly	Ala	Ala	Thr		760 Met	Val	Ala	Ala		765 Leu	Ala	Arg	Ala
		770 Pro	Pro	Leu	Gly		775 Glu	Val	Val	Arg		780 Met	Leu	Asp	Ala	
50	785 Ser	His	Ala	Trp	Ser 805	790 Met	Arg	Lys	Ser	Lys 810	795 Ala	Asn	Trp	Tyr	Arg 815	800 Ile
50	Val	Gly	Val	Leu 820		Trp	Ala	Val	Gly 825		Ala	Lys	Trp	Leu 830		Asn
	Ile	Arg	Arg 835		Arg	Asn	Pro	Val 840		Thr	Val	Leu	Val 845		Ile	Leu
55	Tyr	Leu 850	Val	Leu	Val	Trp	Tyr 855		Asp	Leu	Val	Val 860		Thr	Ala	Phe
	Leu 865		Val	Val	Met	Ile 870	Gly	Val	Trp	Tyr	Tyr 875		Phe	Arg	Pro	Lys 880
60		Pro	Ala	Gly	Met 885	Asp	Ile	Arg	Leu	Ser 890	Gln	Ala	Glu	Thr	Val 895	
	Pro	Asp	Glu	Leu 900	Asp	Glu	Glu	Phe	Asp 905		Ile	Pro	Ser	Ser 910	Arg	Arg

530 Pro Glu Val Ile Arg Ala Arg Tyr Asp Arg Leu Arg Ile Leu Ala Val 920 Arg Val Gln Thr Ile Leu Gly Asp Phe Ala Ala Gln Gly Glu Arg Ile 935 940 5 Gln Ala Leu Val Ser Trp Arg Asp Pro Arg Ala Thr Lys Leu Phe Ile 950 955 Ala Ile Cys Leu Val Ile Thr Ile Val Leu Tyr Ala Val Pro Ala Lys 965 970 Met Val Ala Val Ala Leu Gly Val Ser Asp Ser Val Pro Thr Ala Lvs 10 980 985 990 Gln Asp Thr Lys Glu Ser Leu Lys Lys Ser Phe Ser Ser Leu Arg Phe 995 1000 1005 Asp Phe Ser Ser Met Ala Val Val Gly Ala Pro Ile Ser Ser Pro Ala 1010 1015 1020 15 Ala Gln Leu Gln Thr Gln Phe Leu Ser Asn Pro Ile Leu Pro Arg Phe 1025 1030 1035 1040 Arg Arg Ser Phe Ser Thr Gly Lys Ser Pro Ala Thr Phe Ser Val Val 1045 1050 1055 Ala Met Ala Pro Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys Trp Lys 20 1060 1065 1070 Lys Gln Trp Tyr Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu Gln Ile 1075 1080 1085 Asn Val Asp Val Phe Lys Lys Leu Glu Lys Arg Lys Val Leu Ser Asn 1090 1095 1100 25 Val Glu Lys Ser Gly Leu Leu Ser Lys Ala Glu Gly Leu Gly Leu Thr 1105 1110 1115 1120 Leu Ser Ser Leu Glu Lys Leu Lys Val Phe Ser Lys Ala Glu Asp Leu 1125 1130 Gly Leu Leu Ser Leu Leu Glu Asn Leu Ala Gly Thr Ser Pro Ala Val 30 1140 1145 1150 Leu Ala Ser Ala Ala Leu Pro Ala Leu Thr Ala Ala Ile Val Ala Val 1155 1160 1165 Val Leu Ile Pro Asp Asp Ser Thr Thr Leu Val Val Ala Gln Ala Val 1170 1175 1180 35 Leu Ala Gly Ala Leu Ala Leu Thr Gly Val Val Leu Leu Val Gly Ser 1185 1190 1195 Val Val Leu Asp Gly Leu Gln Glu Ala Asp 1205 (2) INFORMATION FOR SEQ ID NO:403: 40 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1199 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 45 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..1199 (D) OTHER INFORMATION: / Ceres Seq. ID 2028732 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:403: Met Asp Tyr Asp Glu Leu Asp Ile Glu Val Tyr Asn Asp Lys Arg Phe 5 10 Gly Asn Gly Gly Gly Arg Lys Asn His Phe Leu Gly Arg Val Lys Ile 55 Tyr Gly Ser Gln Phe Ser Arg Arg Gly Glu Glu Gly Leu Val Tyr Phe 4.0 Pro Leu Glu Lys Lys Ser Val Phe Ser Trp Ile Arg Gly Glu Ile Gly

Leu Lys Ile Tyr Tyr Tyr Asp Glu Ala Ala Asp Glu Asp Thr Ala Gly

Gly Gly Gly Gln Gln Gln Gln Gln Gln Gln Gln Phe His Pro

90

	Pro	Gln	Gln	Glu 100	Ala	Asp	Glu	Gln		His	Gln	Gln	Gln	Phe 110	His	Pro
	Pro	Pro	Gln 115	Gln	Met	Met	Asn	Ile 120	Pro	Pro	Glu	Lys	Pro 125	Asn	Val	Val
5	Val	Val 130	Glu	Glu	Gly	Arg	Val 135	Phe	Glu	Ser	Ala	Gln 140	Ser	Gln	Arg	Tyr
	Thr 145	Glu	Thr	His	Gln	Gln 150	Pro	Pro	Val	Val	Ile 155	Val	Glu	Glu	Ser	Pro 160
10			His		165					170					175	
	_		His	180		_			185					190		
7 F			Val 195		_	_		200					205			_
15	_	210	Pro	_	_	_	215					220	_			
	225	_	Tyr			230					235		_	_	-	240
20			Met		245	_				250					255	
			Tyr	260			-		265			_	_	270		
٥٢			Ser 275		_		_	280	_				285			-
25		290	Pro				295		_			300	_			
	305		Gln			310		_			315					320
30		_	Ala		325					330					335	
			Gly	340		=		_	345					350	_	-
2 5			Asp 355					360					365			
35	_	370	Asp				375					380				
	385		Ile	_		390					395				_	400
40			Ala		405					410					415	
		_	Leu	420	_				425					430	-	
45			Ala 435					440					445			
40		450	Gln His				455					460				
	465		Glu		_	470					475					480
50			Lys		485					490					495	
			Glu	500					505					510		
55			515 Glu		_		_	520	-				525	_	-	
33		530	Gly	_		_	535	_	_			540				-
	545	_	_	_	_	550					555					560
60			Glu		565					570					575	
	ser	nsp	Phe	580	LIO	T 11T	nid	туз	585	пец	ттÞ	пур	ETO	590	тте	дтХ

```
Ile Leu Glu Leu Gly Ile Leu Gly Ala Arg Gly Leu Leu Pro Met Lys
                ... 600
    Ala Lys Asn Gly Gly Lys Gly Ser Thr Asp Ala Tyr Cys Val Ala Lys
                       615
5
    Tyr Gly Lys Lys Trp Val Arg Thr Arg Thr Ile Thr Asp Ser Phe Asp
               630
    Pro Arg Trp His Glu Gln Tyr Thr Trp Gln Val Tyr Asp Pro Cys Thr
                                 650
    Val Leu Thr Val Gly Val Phe Asp Asn Trp Arg Met Phe Ser Asp Ala
10
                              665
    Ser Asp Asp Arg Pro Asp Thr Arg Ile Gly Lys Ile Arg Ile Arg Val
                680
    Ser Thr Leu Glu Ser Asn Lys Val Tyr Thr Asn Ser Tyr Pro Leu Leu
           695
                                        700
15
    Val Leu Leu Pro Ser Gly Met Lys Lys Met Gly Glu Ile Glu Val Ala
           710 715
    Val Arg Phe Ala Cys Pro Ser Leu Leu Pro Asp Val Cys Ala Ala Tyr
                                 730
                 725
    Gly Gln Pro Leu Leu Pro Arg Met His Tyr Ile Arg Pro Leu Gly Val
20
              740
                              745
    Ala Gln Gln Asp Ala Leu Arg Gly Ala Ala Thr Lys Met Val Ala Ala
           755 760
    Trp Leu Ala Arg Ala Glu Pro Pro Leu Gly Pro Glu Val Val Arg Tyr
       770 775
                                        780
25
    Met Leu Asp Ala Asp Ser His Ala Trp Ser Met Arg Lys Ser Lys Ala
                                     795
    785 790
    Asn Trp Tyr Arg Ile Val Gly Val Leu Ala Trp Ala Val Gly Leu Ala
                 805
                                 810 815
    Lys Trp Leu Asp Asn Ile Arg Arg Trp Arg Asn Pro Val Thr Thr Val
30
                              825
      820
    Leu Val His Ile Leu Tyr Leu Val Leu Val Trp Tyr Pro Asp Leu Val
          835 840 845
     Val Pro Thr Ala Phe Leu Tyr Val Val Met Ile Gly Val Trp Tyr Tyr
       850 855
                            860
35
    Arg Phe Arg Pro Lys Ile Pro Ala Gly Met Asp Ile Arg Leu Ser Gln
                  870 875
     Ala Glu Thr Val Asp Pro Asp Glu Leu Asp Glu Glu Phe Asp Thr Ile
                                 890 895
                 885
     Pro Ser Ser Arg Arg Pro Glu Val Ile Arg Ala Arg Tyr Asp Arg Leu
40
                              905
     Arg Ile Leu Ala Val Arg Val Gln Thr Ile Leu Gly Asp Phe Ala Ala
                           920
     Gln Gly Glu Arq Ile Gln Ala Leu Val Ser Trp Arg Asp Pro Arg Ala
                        935
45
     Thr Lys Leu Phe Ile Ala Ile Cys Leu Val Ile Thr Ile Val Leu Tyr
                     950
                                     955
     Ala Val Pro Ala Lys Met Val Ala Val Ala Leu Gly Val Ser Asp Ser
                                  970
     Val Pro Thr Ala Lys Gln Asp Thr Lys Glu Ser Leu Lys Lys Ser Phe
50
                              985
     Ser Ser Leu Arg Phe Asp Phe Ser Ser Met Ala Val Val Gly Ala Pro
                           1000 1005
     Ile Ser Ser Pro Ala Ala Gln Leu Gln Thr Gln Phe Leu Ser Asn Pro
                       1015
                                        1020
55
     Ile Leu Pro Arg Phe Arg Arg Ser Phe Ser Thr Gly Lys Ser Pro Ala
                                     1035 1040
     1025 1030
     Thr Phe Ser Val Val Ala Met Ala Pro Gln Lys Lys Val Asn Lys Tyr
                 1045
                                  1050 1055
     Asp Ala Lys Trp Lys Lys Gln Trp Tyr Gly Ala Gly Leu Phe Phe Glu
60
             1060 1065 1070
     Gly Ser Glu Gln Ile Asn Val Asp Val Phe Lys Lys Leu Glu Lys Arg
                           1080
                                            1085
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Lys Val Leu Ser Asn Val Glu Lys Ser Gly Leu Leu Ser Lys Ala Glu
                             1095
                                                  1100
     Gly Leu Gly Leu Thr Leu Ser Ser Leu Glu Lys Leu Lys Val Phe Ser
                         1110
                                  1115
 5
     Lys Ala Glu Asp Leu Gly Leu Leu Ser Leu Leu Glu Asn Leu Ala Gly
                                         1130
                     1125
     Thr Ser Pro Ala Val Leu Ala Ser Ala Ala Leu Pro Ala Leu Thr Ala
                                     1145
                 1140
                                                         1150
     Ala Ile Val Ala Val Val Leu Ile Pro Asp Asp Ser Thr Thr Leu Val
10
                                 1160
                                                     1165
     Val Ala Gln Ala Val Leu Ala Gly Ala Leu Ala Leu Thr Gly Val Val
                             1175
                                                 1180
     Leu Leu Val Gly Ser Val Val Leu Asp Gly Leu Gln Glu Ala Asp
                         1190
                                              1195
15
     (2) INFORMATION FOR SEQ ID NO: 404:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 528 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
20
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..528
25
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029079
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:404:
     aaaaaactca ctttcacttg cacaaagata aggaaaccat gtctgtgtca gcgatctttg
                                                                            60
      gtaccggaat cgtcaccgtc gctgcttctc cggttctccg ccaatttcaa gttccaaaat
                                                                            120
      tgggtaatgg aggtggatta gggatggtga ttgagtgttc gtcgaggcca cagaagaaat
                                                                            180
30
     cgacagetea teacaggaag acgaggeega agaagaetea geettgggae attaagagaa
                                                                            240
     agectactgt gtatgeteet etteeteete tteeggegga atggagteeg tttactettg
                                                                            300
     cttccaacga cggtggtgcc gccgttgctg cttctcctgc cggagatttg gtttcaggct
                                                                            360
     ctgcctagtt atatgagcta tctggtggtt tggttgtaat ctagtttgag agcttttgtt
                                                                            420
     ggtgtttgtt tctgttaatg tttttcagtt ttctgctgtc tgtttcttat tatccggttt
35
      atgtttggct ggtgaatgaa tgtcaatgga ccaaactgca atttttt
      (2) INFORMATION FOR SEQ ID NO:405:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 121 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..121
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029080
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:405:
      Lys Thr His Phe His Leu His Lys Asp Lys Glu Thr Met Ser Val Ser
                                          10
                      5
50
      Ala Ile Phe Gly Thr Gly Ile Val Thr Val Ala Ala Ser Pro Val Leu
                                      25
      Arg Gln Phe Gln Val Pro Lys Leu Gly Asn Gly Gly Leu Gly Met
                                  40
      Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser Thr Ala His His
55
                              55
      Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp Ile Lys Arg Lys
                          70
                                              75
      Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala Glu Trp Ser Pro
                                          90
60
      Phe Thr Leu Ala Ser Asn Asp Gly Gly Ala Ala Val Ala Ala Ser Pro
      Ala Gly Asp Leu Val Ser Gly Ser Ala
```

```
115
                                 120
      (2) INFORMATION FOR SEQ ID NO:406:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 109 amino acids
 5
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
10
                (A) NAME/KEY: peptide
               (B) LOCATION: 1..109
               (D) OTHER INFORMATION: / Ceres Seq. ID 2029081
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:406:
     Met Ser Val Ser Ala Ile Phe Gly Thr Gly Ile Val Thr Val Ala Ala
15
                                         10
      Ser Pro Val Leu Arg Gln Phe Gln Val Pro Lys Leu Gly Asn Gly Gly
                                      25
      Gly Leu Gly Met Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser
                                 4.0
20
      Thr Ala His His Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp
                              5.5
      Ile Lys Ard Lys Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala
                          70
      Glu Trp Ser Pro Phe Thr Leu Ala Ser Asn Asp Gly Gly Ala Ala Val
25
                     85
      Ala Ala Ser Pro Ala Gly Asp Leu Val Ser Gly Ser Ala
                  100
      (2) INFORMATION FOR SEQ ID NO:407:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029082
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:407:
40
      Met Val Ile Glu Cys Ser Ser Arg Pro Gln Lys Lys Ser Thr Ala His
                                         10
      His Arg Lys Thr Arg Pro Lys Lys Thr Gln Pro Trp Asp Ile Lys Arg
                                 25
                 20
      Lys Pro Thr Val Tyr Ala Pro Leu Pro Pro Leu Pro Ala Glu Trp Ser
45
                                 40
                                                      45
      Pro Phe Thr Leu Ala Ser Asn Asp Gly Gly Ala Ala Val Ala Ala Ser
                              55
      Pro Ala Gly Asp Leu Val Ser Gly Ser Ala
                         70
50
      (2) INFORMATION FOR SEQ ID NO:408:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 787 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
55
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..787
60
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029225
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:408:
      agggattcta gggctttcct tcattggtca gatcagacga cgttttacat cttcttcttc 60
```

120

180

240

300

420

480

540

600

660

720

780

```
ttcctcttcq atattcqtca qtgtgtgtat tttggggaaa actttgtgag caaaqaqcqa
     gaaaatgagc ggagccggta agaaaatcgc ggatgtggct ttcaaagctt caaggactat
     cgattgggat ggtatggcta aggtccttgt cacagatgag gctcgtagag agttctctaa
     ccttcqtcqt qctttcqatq aggttaacac acagctccag accaaattta qtcaqqaacc
 5
     tgaacctata gattgggatt actataggaa gggtattgga gctggcattg ttgacaagta
     caaqqaaqct tatqacaqca ttqaqattcc aaaqtacqtt qacaaaqtta ctcctqaata
     caaqccaaaq tttqatqctt tqttqqtqqa actqaaaqaa qcaqaacaqa aatcqctcaa
     qqaqtctqaa cqqttqqaqa aaqaaattqc tqatqtccaa qaqatcaqca aaaaqctcaq
     caccatgact gcagatgagt actttgagaa gcaccoggaa ctcaaaaaaga agtttgatga
10
     cqaaatccqt aatqacaact qqqqatactq atcatqtttc tccatctccq qcttqqaaaq
     aaaactctct ttctctttct ctgttctctt actgtgattt tgtgagccaa tcataacaat
     aataagtaca ccattcactt aagcagtgtt gagatcttca ttccaaggaa gataaacgca
     tttggtt
     (2) INFORMATION FOR SEQ ID NO:409:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 168 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..168
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029226
25
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:409:
     Met Ser Gly Ala Gly Lys Lys Ile Ala Asp Val Ala Phe Lys Ala Ser
                                         10
     Arg Thr Ile Asp Trp Asp Gly Met Ala Lys Val Leu Val Thr Asp Glu
                                     2.5
30
     Ala Arg Arg Glu Phe Ser Asn Leu Arg Arg Ala Phe Asp Glu Val Asn
                                 4.0
      Thr Gln Leu Gln Thr Lys Phe Ser Gln Glu Pro Glu Pro Ile Asp Trp
                             55
                                                 60
     Asp Tyr Tyr Arg Lys Gly Ile Gly Ala Gly Ile Val Asp Lys Tyr Lys
35
                         70
                                             75
      Glu Ala Tyr Asp Ser Ile Glu Ile Pro Lys Tyr Val Asp Lys Val Thr
                                         90
                     85
      Pro Glu Tyr Lys Pro Lys Phe Asp Ala Leu Leu Val Glu Leu Lys Glu
                                   105
                 100
40
      Ala Glu Gln Lys Ser Leu Lys Glu Ser Glu Arg Leu Glu Lys Glu Ile
                                 120
                                                     125
      Ala Asp Val Glu Ile Ser Lys Lys Leu Ser Thr Met Thr Ala Asp
                             135
                                                140
      Glu Tyr Phe Glu Lys His Pro Glu Leu Lys Lys Phe Asp Asp Glu
45
                         150
                                            155
      Ile Arg Asn Asp Asn Trp Gly Tyr
                     165
      (2) INFORMATION FOR SEQ ID NO:410:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 145 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..145
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029227
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:410:
60
      Met Ala Lvs Val Leu Val Thr Asp Glu Ala Arg Arg Glu Phe Ser Asn
                                         1.0
      Leu Arg Arg Ala Phe Asp Glu Val Asn Thr Gln Leu Gln Thr Lys Phe
```

536 25 20 Ser Gln Glu Pro Glu Pro Ile Asp Trp Asp Tyr Tyr Arg Lys Gly Ile 40 Gly Ala Gly Ile Val Asp Lys Tyr Lys Glu Ala Tyr Asp Ser Ile Glu 5 55 Ile Pro Lys Tyr Val Asp Lys Val Thr Pro Glu Tyr Lys Pro Lys Phe 75 70 Asp Ala Leu Leu Val Glu Leu Lys Glu Ala Glu Gln Lys Ser Leu Lys 85 90 10 Glu Ser Glu Arg Leu Glu Lys Glu Ile Ala Asp Val Gln Glu Ile Ser 105 Lys Lys Leu Ser Thr Met Thr Ala Asp Glu Tyr Phe Glu Lys His Pro 120 Glu Leu Lys Lys Lys Phe Asp Asp Glu Ile Arg Asn Asp Asn Trp Gly 15 130 135 Tyr 145 (2) INFORMATION FOR SEQ ID NO:411: (i) SEQUENCE CHARACTERISTICS: 20 (A) LENGTH: 1587 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 25 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..1587 (D) OTHER INFORMATION: / Ceres Seq. ID 2029422 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:411: 30 gtagtototo tttttataac cacttotoga aaactgaaac ctttgtagag agaacccata 60 gttcgataaa acattctttt tgcaactgag acttggcaac ttggttttac tcaaagtaag 120 atttcttctt gttgatggtg ataagttcaa tatgatgaaa agtttggcta gtgcggttgg agggaagacg gcgagggcat gtgatagctg cgtgaagagg cgggcacgtt ggtattgcgc 240 agctgatgat gcctttcttt gccatgcttg tgacggttcg gtccactcgg caaaccctct 300 35 360 tgctcgtagg cacgagagag ttcgcttgaa atcggctagc gccggaaagt atcgccatgc ctcgccgcct caccaagcca cgtggcatca gggatttaca Cgtaaagctc ggaccccacg 420 tggaggcaag aagagccaca cgatggtttt tcatgatctt gtgccggaga tgagcacgga 480 540 ggatcaagog gagagttaog aggtggaaga goagotcata titgaggtgo oggtgatgaa ctcgatggtt qaqqaqcaat gctttaacca atccctggag aaacagaatg agtttccaat 600 40 gatgccctta aqtttcaaqa qtagtgacga agaagatgac gacaacgctg agagttgtct 660 gaatggtttg ttcccaaccg acatggaact agctcagttc acagctgacg tggagactct 720 actcggtgga ggggatcgag agtttcattc catagaagaa ctagggttag gtgagatgtt 780 840 aaaqatcqaa aaaqaqgaqq tggaggaaga gggagttgtg acaagagaag tgcatgatca agatgaaggt gatgagacat ccccatttga aataagcttt gactacgagt acacacacaa 900 45 960 gaccacattc gatgaaggag aagaagatga gaaagaagac gtgatgaaga atgtgatgga gatgggagtg aatgagatga gtggtgggat taaagaagag aagaaggaga aggctcttat 1020 gcttagattg gactatgaat cagtcatttc cacttgggga ggccaaggga tcccatggac 1080 cgcccgggtg ccatctgaaa tagacctcga catggtttgt ttcccaaccc ataccatggg 1140 tgaaagtgga qcagaggctc atcatcacaa ccacttccgc ggcctagggt tacacctagg 1200 50 agatgctqqq qatqqaqqaa gagaggctag ggtttcaaga taccqagaga aaaggaggac aaggttgttc tccaaqaaqa taaggtacqa qgtacgtaaa ttgaatgcag agaaaaggcc tcgcatgaaa ggaaggttcg tcaagagatc ttcaattggt gttgctcact aaagaactta attaattatq qatattaaat tactttqctc tcatcttqct tttttqttqc tataqttttq qtqattqtta qctttctttt tctqcattca taqaqaattt tqcacqtttt tqtqaqctac 55 qtatqtacat aaatatatca ccaaaaaatq tqactatctt qtaaqcactq atttatataq togatataac gtgaattttg attgctg (2) INFORMATION FOR SEQ ID NO:412: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 406 amino acids 60 (B) TYPE: amino acid (C) STRANDEDNESS:

(D) TOPOLOGY: linear

537 (ii) MOLECULE TYPE: peptide

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(ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..406
5
             (D) OTHER INFORMATION: / Ceres Seq. ID 2029423
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:412:
     Met Met Lys Ser Leu Ala Ser Ala Val Gly Gly Lys Thr Ala Arg Ala
                                   10
     Cys Asp Ser Cys Val Lys Arg Arg Ala Arg Trp Tyr Cys Ala Ala Asp
10
         20
                             25
     Asp Ala Phe Leu Cys His Ala Cys Asp Gly Ser Val His Ser Ala Asn
                             40
     Pro Leu Ala Arg Arg His Glu Arg Val Arg Leu Lys Ser Ala Ser Ala
                         55
15
     Gly Lys Tyr Arg His Ala Ser Pro Pro His Gln Ala Thr Trp His Gln
                                       75
     Gly Phe Thr Arg Lys Ala Arg Thr Pro Arg Gly Gly Lys Lys Ser His
     Thr Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln
20
                                105
               100
     Ala Glu Ser Tyr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val
                            120
                                              125
           115
     Met Asn Ser Met Val Glu Glu Glr Cvs Phe Asn Gln Ser Leu Glu Lys
                      135
                                        1.40
25
     Gln Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu
         150 155
     Glu Asp Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr
                  165
                                  170
     Asp Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly
30
               180
                   185
     Gly Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu
                             200
                                               205
     Met Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Gly Val Val Thr
                         215
35
     Arg Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe Glu
                     230
                                      235
     Ile Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu Gly
                                   250
                   245
     Glu Glu Asp Glu Lys Glu Asp Val Met Lys Asn Val Met Glu Met Gly
40
                                265
               260
     Val Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys Ala
                            280
                                              285
     Leu Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly Gly
                         295
                                          300
45
     Gln Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu Asp
                                       315
         310
     Met Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu Ala
                                    330
                  325
     His His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp Ala
50
              340 345 350
     Gly Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys Arg
           355
                360
     Arg Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys Leu
                         375 380
55
     Asn Ala Glu Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg Ser
                     390
                                       395
     Ser Ile Gly Val Ala His
                  405
     (2) INFORMATION FOR SEQ ID NO:413:
60
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 405 amino acids
              (B) TYPE: amino acid
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538

(C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 5 (A) NAME/KEY: peptide (B) LOCATION: 1..405 (D) OTHER INFORMATION: / Ceres Seq. ID 2029424 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:413: Met Lys Ser Leu Ala Ser Ala Val Gly Gly Lys Thr Ala Arg Ala Cys 10 10 Asp Ser Cys Val Lys Arg Arg Ala Arg Trp Tyr Cys Ala Ala Asp Asp 25 Ala Phe Leu Cys His Ala Cys Asp Gly Ser Val His Ser Ala Asn Pro 40 15 Leu Ala Arg Arg His Glu Arg Val Arg Leu Lys Ser Ala Ser Ala Gly 55 60 Lys Tyr Arg His Ala Ser Pro Pro His Gln Ala Thr Trp His Gln Glv 70 7.5 Phe Thr Arg Lys Ala Arg Thr Pro Arg Gly Gly Lys Lys Ser His Thr 20 90 Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln Ala 105 Glu Ser Tyr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val Met 120 115 25 Asn Ser Met Val Glu Glu Gln Cys Phe Asn Gln Ser Leu Glu Lys Gln 135 140 Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu Glu 150 155 Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr Asp 30 165 170 Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly Gly 180 185 Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu Met 195 200 205 35 Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Gly Val Val Thr Arg 215 220 Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe Glu Ile 230 235 Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu Gly Glu 40 245 250 Glu Asp Glu Lys Glu Asp Val Met Lys Asn Val Met Glu Met Gly Val 265 Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys Ala Leu 280 45 Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly Gly Gln 300 295 Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu Asp Met 310 315 Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu Ala His 325 330 50 His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp Ala Gly 345 Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys Arg Arg 360 355 55 Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys Leu Asn 375 380 Ala Glu Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg Ser Ser 395 Ile Gly Val Ala His 60 405

(2) INFORMATION FOR SEQ ID NO:414: (i) SEQUENCE CHARACTERISTICS:

```
(A) LENGTH: 309 amino acids
             (B) TYPE: amino acid
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
5
       (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..309
             (D) OTHER INFORMATION: / Ceres Seq. ID 2029425
10
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:414:
     Met Val Phe His Asp Leu Val Pro Glu Met Ser Thr Glu Asp Gln Ala
     Glu Ser Tvr Glu Val Glu Glu Gln Leu Ile Phe Glu Val Pro Val Met
                   25
     Asn Ser Met Val Glu Glu Gln Cys Phe Asn Gln Ser Leu Glu Lys Gln
15
                  40
     Asn Glu Phe Pro Met Met Pro Leu Ser Phe Lys Ser Ser Asp Glu Glu
                      55
     Asp Asp Asp Asn Ala Glu Ser Cys Leu Asn Gly Leu Phe Pro Thr Asp
20
                   70
                              75 80
     Met Glu Leu Ala Gln Phe Thr Ala Asp Val Glu Thr Leu Leu Gly Gly
                 85 90 95
     Gly Asp Arg Glu Phe His Ser Ile Glu Glu Leu Gly Leu Gly Glu Met
              100 105 110
     Leu Lys Ile Glu Lys Glu Glu Val Glu Glu Glu Gly Val Val Thr Arg
25
           115 120 125
     Glu Val His Asp Gln Asp Glu Gly Asp Glu Thr Ser Pro Phe Glu Ile
       130 135
     Ser Phe Asp Tyr Glu Tyr Thr His Lys Thr Thr Phe Asp Glu Gly Glu
30
                                       155
                     150
     Glu Asp Glu Lys Glu Asp Val Met Lys Asn Val Met Glu Met Gly Val
                                   170
                  165
     Asn Glu Met Ser Gly Gly Ile Lys Glu Glu Lys Lys Glu Lys Ala Leu
                                185 190
35
     Met Leu Arg Leu Asp Tyr Glu Ser Val Ile Ser Thr Trp Gly Gly Gln
                            200
     Gly Ile Pro Trp Thr Ala Arg Val Pro Ser Glu Ile Asp Leu Asp Met
                        215
     Val Cys Phe Pro Thr His Thr Met Gly Glu Ser Gly Ala Glu Ala His
40
                     230
                                       235
     His His Asn His Phe Arg Gly Leu Gly Leu His Leu Gly Asp Ala Gly
                                    250
                  245
     Asp Gly Gly Arg Glu Ala Arg Val Ser Arg Tyr Arg Glu Lys Arg Arg
                                265
     Thr Arg Leu Phe Ser Lys Lys Ile Arg Tyr Glu Val Arg Lys Leu Asn
4.5
           275 280 285
     Ala Glu Lys Arg Pro Arg Met Lys Gly Arg Phe Val Lys Arg Ser Ser
                         295
       290
     Ile Gly Val Ala His
50
     305
     (2) INFORMATION FOR SEQ ID NO:415:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 1300 base pairs
             (B) TYPE: nucleic acid
55
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: DNA (genomic)
         (ix) FEATURE:
             (A) NAME/KEY: -
60
             (B) LOCATION: 1..1300
             (D) OTHER INFORMATION: / Ceres Seq. ID 2029806
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:415:
```

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ctcaaqtttc ttqcaattac ctacttacaa caaaqcaact cqactcqaac aaacacatcc
     atggeteett etgegeaace tetteetgtg agtgtttegg atgaaaaata tgegaatgte
                                                                          120
     aagtgggaag agttggcatt caagtttgtt cgtacggatt atatgtatgt tgcgaagtgc
                                                                          180
     aatcatggag agagttttca agaggggaag attcttcctt ttgctgattt gcaacttaac
                                                                          240
 5
     ccttgcgctg ctgttcttca gtatggccag ggtttatatg aaggactgaa agcttacagg
                                                                          300
     acagaagatg gtcggattct gctattccga ccagaccaaa acggtctccg ccttcaagcc
                                                                          360
     ggagetgaca gactetatat gcettateet teggtegate aattegtete egecateaaa
                                                                          420
     caagttgctc ttgccaacaa gaaatggatt cctcctccgg ggaaaggaac attgtatatt
                                                                           480
     aggectatet tgtttgggag tggtccgatt cttggttcat ttcccattcc tgagaccacc
                                                                           540
10
     ttcacagctt ttgcctgtcc tgttggacgt tatcataagg ataactctgg tttgaatctg
                                                                           600
     aaaatcgaag atcagtttcg tcgagctttt cctagtggaa ctggtggtgt gaaqaqcatc
                                                                           660
     acaaactatt gtcctgtttg gataccattg gcagaggcga aaaaacaagg tttctctgat
     attttqtttt tqqatqctqc aactggcaaa aacattgaag aacttttcgc agctaatqtt
                                                                          780
     tttatgctca agggcaatgt tgtatcgaca ccaacaattg caggaactat tttgcccgga
                                                                          840
15
     gtcactcgaa actgcgtaat ggaattgtgt cgtgatttcg gctaccaggt cgaggaacgt
                                                                          900
     acgattcctc tagtggactt tctcgatqcg gacgaagctt tctgtactgg cactgcttcc
                                                                          960
     attgtgacta gtattgcatc cgtaaccttt aaagacaaaa agaccggatt caaaacaggg
                                                                         1020
     1080
     gaggatacca agggatggac ggtggagatt gaccgccagg gctgaaagtt gaaactgtaa
                                                                         1140
20
     cttgatgcta aatatgtgtg tgtgtgtata caaaacttat gtaagaaaca tctgaagatg
                                                                         1200
     tototgatot tigtgatigt gatgatoatg catgitottat cgitgcgatg tattitataa
                                                                         1260
     atgttcgtct gtaagttatt taataactat ggctttttgc
      (2) INFORMATION FOR SEQ ID NO:416:
          (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 374 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
30
         (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..374
                (D) OTHER INFORMATION: / Ceres Seq. ID 2029807
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:416:
35
     Leu Lys Phe Leu Ala Ile Thr Tyr Leu Gln Gln Ser Asn Ser Thr Arg
                                         10
     Thr Asn Thr Ser Met Ala Pro Ser Ala Gln Pro Leu Pro Val Ser Val
                                     25
                 2.0
      Ser Asp Glu Lys Tyr Ala Asn Val Lys Trp Glu Glu Leu Ala Phe Lys
40
                                40
     Phe Val Arg Thr Asp Tyr Met Tyr Val Ala Lys Cys Asn His Gly Glu
                             55
     Ser Phe Gln Glu Gly Lys Ile Leu Pro Phe Ala Asp Leu Gln Leu Asn
                         70
                                             7.5
45
      Pro Cys Ala Ala Val Leu Gln Tyr Gly Gln Gly Leu Tyr Glu Gly Leu
                                         90
                     8.5
     Lys Ala Tyr Arg Thr Glu Asp Gly Arg Ile Leu Leu Phe Arg Pro Asp
                                     105
                 100
                                                         110
     Gln Asn Gly Leu Arg Leu Gln Ala Gly Ala Asp Arg Leu Tyr Met Pro
50
                                                     125
             115
                                 120
     Tyr Pro Ser Val Asp Gln Phe Val Ser Ala Ile Lys Gln Val Ala Leu
                             135
                                                 140
     Ala Asn Lys Lys Trp Ile Pro Pro Pro Gly Lys Gly Thr Leu Tyr Ile
                         150
                                             155
55
     Arg Pro Ile Leu Phe Gly Ser Gly Pro Ile Leu Gly Ser Phe Pro Ile
                     165
                                         170
      Pro Glu Thr Thr Phe Thr Ala Phe Ala Cys Pro Val Gly Arg Tyr His
                 180
                                     185
     Lys Asp Asn Ser Gly Leu Asn Leu Lys Ile Glu Asp Gln Phe Arg Arg
60
                                200
          195
     Ala Phe Pro Ser Gly Thr Gly Gly Val Lys Ser Ile Thr Asn Tyr Cys
                             215
          210
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Pro Val Trp Ile Pro Leu Ala Glu Ala Lys Lys Gln Gly Phe Ser Asp
                                  235
    225 230
    Ile Leu Phe Leu Asp Ala Ala Thr Gly Lys Asn Ile Glu Glu Leu Phe
           245 250 255
    Ala Ala Asn Val Phe Met Leu Lys Gly Asn Val Val Ser Thr Pro Thr
            260 265 270
    Ile Ala Gly Thr Ile Leu Pro Gly Val Thr Arg Asn Cys Val Met Glu
     275 280 285
    Leu Cys Arg Asp Phe Gly Tyr Gln Val Glu Glu Arg Thr Ile Pro Leu
10
     290 295 300
    Val Asp Phe Leu Asp Ala Asp Glu Ala Phe Cys Thr Gly Thr Ala Ser
    305 310 315
    Ile Val Thr Ser Ile Ala Ser Val Thr Phe Lys Asp Lys Lys Thr Gly
                325 330 335
15
    Phe Lys Thr Gly Glu Glu Thr Leu Ala Ala Lys Leu Tyr Glu Thr Leu
          340 345 350
    Ser Asp Ile Gln Thr Gly Arg Val Glu Asp Thr Lys Gly Trp Thr Val
         355 360
    Glu Ile Asp Arg Gln Gly
20
      370
    (2) INFORMATION FOR SEQ ID NO:417:
        i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 354 amino acids
            (B) TYPE: amino acid
25
           (C) STRANDEDNESS:
            (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: peptide
       (ix) FEATURE:
           (A) NAME/KEY: peptide
30
           (B) LOCATION: 1..354
           (D) OTHER INFORMATION: / Ceres Seq. ID 2029808
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:417:
    Met Ala Pro Ser Ala Gln Pro Leu Pro Val Ser Val Ser Asp Glu Lys
       5 10
35
    Tyr Ala Asn Val Lys Trp Glu Glu Leu Ala Phe Lys Phe Val Arg Thr
                    25
    Asp Tyr Met Tyr Val Ala Lys Cys Asn His Gly Glu Ser Phe Gln Glu
                    40
    Gly Lws Ile Leu Pro Phe Ala Asp Leu Gln Leu Asn Pro Cvs Ala Ala
40
     50 55
    Val Leu Gln Tyr Gly Gln Gly Leu Tyr Glu Gly Leu Lys Ala Tyr Arg
                               75 80
                  70
    Thr Glu Asp Gly Arg Ile Leu Leu Phe Arg Pro Asp Gln Asn Gly Leu
                85 90 95
45
    Arg Leu Gln Ala Gly Ala Asp Arg Leu Tyr Met Pro Tyr Pro Ser Val
                           105 110
    Asp Gin Phe Val Ser Ala Ile Lys Gln Val Ala Leu Ala Asn Lys Lys
                         120 125
    Trp Ile Pro Pro Pro Gly Lys Gly Thr Leu Tyr Ile Arg Pro Ile Leu
50
     130 135 140
    Phe Gly Ser Gly Pro Ile Leu Gly Ser Phe Pro Ile Pro Glu Thr Thr
    145 150 155 160
    Phe Thr Ala Phe Ala Cys Pro Val Gly Arg Tyr His Lys Asp Asn Ser
               165 170 175
55
    Gly Leu Asn Leu Lys Ile Glu Asp Gln Phe Arg Arg Ala Phe Pro Ser
            180 185 190
    Gly Thr Gly Gly Val Lys Ser Ile Thr Asn Tyr Cys Pro Val Trp Ile
         195 200 205
    Pro Leu Ala Glu Ala Lys Lys Gln Gly Phe Ser Asp Ile Leu Phe Leu
60
                    215 220
    Asp Ala Ala Thr Gly Lys Asn Ile Glu Glu Leu Phe Ala Ala Asn Val
                   230
                                  235
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542 Phe Met Leu Lys Gly Asn Val Val Ser Thr Pro Thr Ile Ala Gly Thr 250 245 Ile Leu Pro Gly Val Thr Arg Asn Cys Val Met Glu Leu Cys Arg Asp 260 265 270 5 Phe Gly Tyr Gln Val Glu Glu Arg Thr Ile Pro Leu Val Asp Phe Leu 275 280 285 Asp Ala Asp Glu Ala Phe Cys Thr Gly Thr Ala Ser Ile Val Thr Ser 290 295 300 Ile Ala Ser Val Thr Phe Lys Asp Lys Lys Thr Gly Phe Lys Thr Gly 10 310 315 Glu Glu Thr Leu Ala Ala Lys Leu Tyr Glu Thr Leu Ser Asp Ile Gln 325 330 Thr Gly Arg Val Glu Asp Thr Lys Gly Trp Thr Val Glu Ile Asp Arg 345 15 Gln Gly (2) INFORMATION FOR SEQ ID NO:418: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 320 amino acids 20 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 25 (A) NAME/KEY: peptide (B) LOCATION: 1..320 (D) OTHER INFORMATION: / Ceres Seq. ID 2029809 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:418: Met Tyr Val Ala Lys Cys Asn His Gly Glu Ser Phe Gln Glu Gly Lys 10 30 Ile Leu Pro Phe Ala Asp Leu Gln Leu Asn Pro Cys Ala Ala Val Leu 20 25 Gln Tyr Gly Gln Gly Leu Tyr Glu Gly Leu Lys Ala Tyr Arg Thr Glu 40 35 Asp Gly Arg Ile Leu Leu Phe Arg Pro Asp Gln Asn Gly Leu Arg Leu 55 Gln Ala Gly Ala Asp Arg Leu Tyr Met Pro Tyr Pro Ser Val Asp Gln 70 75 Phe Val Ser Ala Ile Lys Gln Val Ala Leu Ala Asn Lys Lys Trp Ile 40 85 90 95 Pro Pro Pro Gly Lys Gly Thr Leu Tyr Ile Arg Pro Ile Leu Phe Gly 100 105 110 Ser Gly Pro Ile Leu Gly Ser Phe Pro Ile Pro Glu Thr Thr Phe Thr 115 120 125 45 Ala Phe Ala Cys Pro Val Gly Arg Tyr His Lys Asp Asn Ser Gly Leu 130 135 140 Asn Leu Lys Ile Glu Asp Gln Phe Arg Arg Ala Phe Pro Ser Gly Thr 150 155 160 Gly Gly Val Lys Ser Ile Thr Asn Tyr Cys Pro Val Trp Ile Pro Leu 50 165 170 175 Ala Glu Ala Lys Lys Gln Gly Phe Ser Asp Ile Leu Phe Leu Asp Ala 180 185 190 Ala Thr Gly Lys Asn Ile Glu Glu Leu Phe Ala Ala Asn Val Phe Met 195 200 55 Leu Lys Glv Asn Val Val Ser Thr Pro Thr Ile Ala Gly Thr Ile Leu 215 220 Pro Gly Val Thr Arg Asn Cys Val Met Glu Leu Cys Arg Asp Phe Gly 230 235 240 Tyr Gln Val Glu Glu Arg Thr Ile Pro Leu Val Asp Phe Leu Asp Ala 60 250 245 Asp Glu Ala Phe Cys Thr Gly Thr Ala Ser Ile Val Thr Ser Ile Ala 265

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543
Ser Val Thr Phe Lys Asp Lys Lys Thr Gly Phe Lys Thr Gly Glu Glu
                            280
Thr Leu Ala Ala Lys Leu Tyr Glu Thr Leu Ser Asp Ile Gln Thr Gly
                        295
Arg Val Glu Asp Thr Lys Gly Trp Thr Val Glu Ile Asp Arg Gln Gly
                    310
                                        315
(2) INFORMATION FOR SEQ ID NO:419:
     (i) SEQUENCE CHARACTERISTICS:
          (A) LENGTH: 987 base pairs
          (B) TYPE: nucleic acid
          (C) STRANDEDNESS: single
          (D) TOPOLOGY: linear
    (ii) MOLECULE TYPE: DNA (genomic)
    (ix) FEATURE:
          (A) NAME/KEY: -
          (B) LOCATION: 1..987
          (D) OTHER INFORMATION: / Ceres Seq. ID 2030038
    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:419:
atatotgaaa gaotoaaaaa ootatogtoa tttatottoa acaggoggaa taacggagat
                                                                        60
ggccgccatt acagetetea eteteogete teetgtttat eteccateat etgecacaag
                                                                       120
soctagatte catggettea ceaatsaace accaecaget egietetiet tissistaa
                                                                       180
spectroset tetetateaa teeaaaacee taaateeate sgaattteeg cateegette
                                                                       240
googataaca acaccaatco tocaqacqqa qaagtcaaca gotoqqtcat cdacactcac
                                                                       300
egggteeact eggteteteg etactetege ggetttagea ategetgtaa eeagagteet
                                                                       360
ageteagaaa eteteteteg caateeagae tteaagteee gtaategeeg aegggttaeg
                                                                       420
attetetete agtacegeeg gacetgtett ettegegtet eteegggate gteeteeggg
                                                                       480
atacttgaac acgccgctga cggttgttgc ggtggggata aagaagtggc tagacattta
                                                                       540
cagtggggta ttgatggtta gggttttgct gagttggttc cctaatatcc cttgggaaag
acagoetttg tetgecatta gagatetetg tgateettae ttgaatetet teagaaacat
                                                                       660
cattcctcct atcttcgata cgcttgatgt tagtccattg cttgctttcg cggttcttgg
                                                                       720
tacacttgga tcgattgttc atggcagcac tgggtagaaa ttggaagact tgagctaatg
gttttgttga aacgttcaat tgggaaattt ctgcaacttg tttcccaaga ttagggaaaa
                                                                       840
atcagaattt gatgctgtgt tggatgctcg atgagagttg tgagctt at cgtttttgtt
                                                                       900
gttctctttt tgactttgat tctgaataat gagattttgg gttgttgtt. atctcctaat
tacatgttca gactaccgtt tattgat
(2) INFORMATION FOR SEQ ID NO:420:
     (i) SEQUENCE CHARACTERISTICS:
          (A) LENGTH: 251 amino acids
          (B) TYPE: amino acid
          (C) STRANDEDNESS:
          (D) TOPOLOGY: linear
   (ii) MOLECULE TYPE: peptide
    (ix) FEATURE:
          (A) NAME/KEY: peptide
          (B) LOCATION: 1..251
          (D) OTHER INFORMATION: / Ceres Seq. ID 2030039
    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:420:
Tyr Leu Lys Asp Ser Lys Thr Tyr Arg His Leu Ser Ser Thr Gly Gly
                                    10
Ile Thr Glu Met Ala Ala Ile Thr Ala Leu Thr Leu Arg Ser Pro Val
                                25
Tyr Leu Pro Ser Ser Ala Thr Ser Pro Arg Phe His Gly Phe Thr Asn
                            40
Gln Pro Pro Pro Ala Arg Leu Phe Phe Pro Leu Asn Pro Phe Pro Ser
Leu Ser Ile Gln Asn Pro Lys Ser Ile Arg Ile Ser Ala Ser Ala Ser
                    7.0
                                        75
Pro Ile Thr Thr Pro Ile Leu Gln Thr Glu Lys Ser Thr Ala Arg Ser
                                    90
Ser Thr Leu Thr Gly Ser Thr Arg Ser Leu Ala Thr Leu Ala Ala Leu
```

\$100\$ \$105\$ \$110\$ Ala Ile Ala Val Thr Arg Val Leu Ala Gln Lys Leu Ser Leu Ala Ile

544 120 Gln Thr Ser Ser Pro Val Ile Ala Asp Gly Leu Arg Phe Ser Leu Ser 135 140 Thr Ala Gly Pro Val Phe Phe Ala Ser Leu Arg Asp Arg Pro Pro Gly 150 5 155 Tyr Leu Asn Thr Pro Leu Thr Val Val Ala Val Gly Ile Lys Lys Trp 165 170 175 Leu Asp Ile Tyr Ser Gly Val Leu Met Val Arg Val Leu Leu Ser Trp 180 185 10 Phe Pro Asn Ile Pro Trp Glu Arg Gln Pro Leu Ser Ala Ile Arg Asp 200 205 Leu Cys Asp Pro Tyr Leu Asn Leu Phe Arg Asn Ile Ile Pro Pro Ile 210 215 220 Phe Asp Thr Leu Asp Val Ser Pro Leu Leu Ala Phe Ala Val Leu Gly 15 225 230 235 Thr Leu Gly Ser Ile Val His Gly Ser Thr Gly 245 (2) INFORMATION FOR SEQ ID NO:421: (i) SEQUENCE CHARACTERISTICS: 20 (A) LENGTH: 232 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 25 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..232 (D) OTHER INFORMATION: / Ceres Seq. ID 2030040 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:421: 30 Met Ala Ala Ile Thr Ala Leu Thr Leu Arg Ser Pro Val Tyr Leu Pro 5 10 Ser Ser Ala Thr Ser Pro Arg Phe His Gly Phe Thr Asn Gln Pro Pro 25 Pro Ala Arg Leu Phe Phe Pro Leu Asn Pro Phe Pro Ser Leu Ser Ile 35 40 Gln Asn Pro Lys Ser Ile Arg Ile Ser Ala Ser Ala Ser Pro Ile Thr 55 Thr Pro Ile Leu Gln Thr Glu Lys Ser Thr Ala Arg Ser Ser Thr Leu 70 75 40 Thr Gly Ser Thr Arg Ser Leu Ala Thr Leu Ala Ala Leu Ala Ile Ala 90 8.5 Val Thr Arg Val Leu Ala Gln Lys Leu Ser Leu Ala Ile Gln Thr Ser 105 Ser Pro Val Ile Ala Asp Gly Leu Arg Phe Ser Leu Ser Thr Ala Gly 4.5 115 120 Pro Val Phe Phe Ala Ser Leu Arg Asp Arg Pro Pro Gly Tyr Leu Asn 135 Thr Pro Leu Thr Val Val Ala Val Gly Ile Lys Lys Trp Leu Asp Ile 150 155 50 Tyr Ser Gly Val Leu Met Val Arg Val Leu Leu Ser Trp Phe Pro Asn 165 170 Ile Pro Trp Glu Arg Gln Pro Leu Ser Ala Ile Arg Asp Leu Cys Asp 185 Pro Tyr Leu Asn Leu Phe Arg Asn Ile Ile Pro Pro Ile Phe Asp Thr 55 195 200 Leu Asp Val Ser Pro Leu Leu Ala Phe Ala Val Leu Gly Thr Leu Gly 215 220 Ser Ile Val His Gly Ser Thr Gly

(2) INFORMATION FOR SEQ ID NO:422: 60 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 755 base pairs

```
(B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
 5
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..755
                (D) OTHER INFORMATION: / Ceres Seq. ID 2031778
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:422:
10
      ccgcagaaaa ccctattgct cccctttgct caagctcagt gcctctttgc agcgaaagct
                                                                              60
      caaccaacaa gacgatgatt atctcagaga acaatcgcag agagatctgc aaataccttt
                                                                             120
      tcaaagaggg agtatgcttt gctaagaagg atttcaatct cccaaagcat ccgttgattg
                                                                             180
      atgtaccaaa cctgcaagtg attaagctca tgcagagttt caagtccaag gagtacgtta
                                                                             240
      gggagacatt tgcgtggatg cattattatt ggtttctgac taatgaagga attgagttct
                                                                             300
15
      tgagaactta tcttaacctt ccttccgatg ttgtccctgc tactttgaag aagtctgcta
                                                                             360
      agcccggtgg tcgtcccttt ggtggcccac ctggtgatcg ccaaagagga ccacctcgct
                                                                             420
      ctgatggaga ccgtcccaga tttggtgacc gtgatggata ccgtggaggc ccacgtggtg
                                                                             480
      gtgatgagaa gggtggagct ccagctgatt tccagccgtc tttccaagga ggtggtggta
                                                                             540
      ggcctggttt tggccgtggt gcaggcggtt acagtgcagc agcaccatct ggttcagggt
                                                                             600
20
                                                                             660
      tcccttgaaa aatttgttgt catattgcga cgatggaagg acagttttgt tttttgttct
                                                                             720
      agttetgttt gtgtaatgea aatetggaat etataateta tetattaeet teacettggt
      ttaaacgaac aaatcctgta gtttcaggat tagtg
      (2) INFORMATION FOR SEQ ID NO:423:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 201 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..201
                (D) OTHER INFORMATION: / Ceres Seq. ID 2031779
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:423:
35
     Ala Glu Asn Pro Ile Ala Pro Leu Cys Ser Ser Ser Val Pro Leu Cys
                                          10
      Ser Glu Ser Ser Thr Asn Lys Thr Met Ile Ile Ser Glu Asn Asn Arg
                                      25
      Arg Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly Val Cys Phe Ala Lys
40
                                  40
      Lys Asp Phe Asn Leu Pro Lys His Pro Leu Ile Asp Val Pro Asn Leu
                              55
      Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg
                                              75
                          70
45
      Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly
                                           90
                      85
      Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro Ser Asp Val Val Pro
                  100
                                      105
      Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly
50
              115
                                  120
      Pro Pro Gly Asp Arg Gln Arg Gly Pro Pro Arg Ser Asp Gly Asp Arg
                              135
                                                   140
      Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Gly Gly Pro Arg Gly Gly
                                               155
                          150
55
      Asp Glu Lys Gly Gly Ala Pro Ala Asp Phe Gln Pro Ser Phe Gln Gly
                      165
                                          170
      Gly Gly Gly Arg Pro Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala
                  180
                                      185
      Ala Ala Pro Ser Gly Ser Gly Phe Pro
60
              195
                                  200
      (2) INFORMATION FOR SEQ ID NO: 424:
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(i) SEQUENCE CHARACTERISTICS:

```
546
               (A) LENGTH: 177 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
 5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..177
               (D) OTHER INFORMATION: / Ceres Seq. ID 2031780
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:424:
     Met Ile Ile Ser Glu Asn Asn Arg Glu Ile Cys Lys Tyr Leu Phe
                                       10
     Lys Glu Gly Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Pro Lys His
                 20
                        25
15
     Pro Leu Ile Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser
                        40
     Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr
                    55
     Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu
20
                        70
                                           75
     Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys
                                       90
     Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Gln Arg Gly
                                   105
25
     Pro Pro Arg Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly
                               120
     Tyr Arg Gly Gly Pro Arg Gly Gly Asp Glu Lys Gly Gly Ala Pro Ala
                           135
     Asp Phe Gln Pro Ser Phe Gln Gly Gly Gly Arg Pro Gly Phe Gly
30
     145 150
                                          155
     Arg Gly Ala Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser Gly Phe
                                      170
                    165
     Pro
35
     (2) INFORMATION FOR SEQ ID NO: 425:
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 132 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
40
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..132
45
               (D) OTHER INFORMATION: / Ceres Seq. ID 2031781
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:425:
     Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp
                                       10
     Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg
50
                 20
                                    25
     Thr Tyr Leu Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys
                                4.0
     Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg
55
     Gln Arg Gly Pro Pro Arg Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp
                        70
                                           7.5
     Arg Asp Gly Tyr Arg Gly Gly Pro Arg Gly Gly Asp Glu Lys Gly Gly
                                       90
     Ala Pro Ala Asp Phe Gln Pro Ser Phe Gln Gly Gly Gly Arg Pro
                     105
60
                100
```

Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Ala Pro Ser Gly
115 120 125

-

```
547
      Ser Gly Phe Pro
          130
      (2) INFORMATION FOR SEQ ID NO: 426:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 538 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
10
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..538
                (D) OTHER INFORMATION: / Ceres Seq. ID 2032723
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 426:
15
      aaagcaataa aaaacaaaaa tggcgtttac tccgaagatc atcacatgcc tcattgtcct
                                                                             60
      tacqatctac atqqcatccc caacagagtc aaccatccag tgtgggacag tgacqagcac
                                                                             120
      actqqcacaq tqcctqacct acttqaccaa caqtqqtcca ttqccatcac aatqctqcqt
                                                                            180
      gggagtcaag tcattgtacc aattggctca gaccacccg gaccgtaaac aagtatgtga
      qtqccttaaa ctaqcqqqta aaqaaatcaa qqqcctcaac accqaccttq tqqccqcact
20
      toctaccact tgtqqtqttt caattoocta coccatcagt tttagcacca attgcqacaq
      tatategact geogtgtgaa agaggetagt gateagatgt aegactaate aaacttgeea
                                                                            420
      gottttaacc taattaaata aaagtattot gottatattt cocattttat gattttatot
                                                                            480
      tottatotat qtaaccacac qatttcatat gctaataatg acaacggato tttototo
      (2) INFORMATION FOR SEQ ID NO: 427:
25
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 119 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..119
                (D) OTHER INFORMATION: / Ceres Seq. ID 2032724
35
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:427:
      Met Ala Phe Thr Pro Lys Ile Ile Thr Cys Leu Ile Val Leu Thr Ile
                                           10
      Tyr Met Ala Ser Pro Thr Glu Ser Thr Ile Gln Cys Gly Thr Val Thr
                  20
                                       25
40
      Ser Thr Leu Ala Gln Cys Leu Thr Tyr Leu Thr Asn Ser Gly Pro Leu
                                  40
      Pro Ser Gln Cys Cys Val Gly Val Lys Ser Leu Tyr Gln Leu Ala Gln
                              55
      Thr Thr Pro Asp Arg Lys Gln Val Cys Glu Cys Leu Lys Leu Ala Gly
45
      Lys Glu Ile Lys Gly Leu Asn Thr Asp Leu Val Ala Ala Leu Pro Thr
                                          90
                      8.5
      Thr Cys Gly Val Ser Ile Pro Tyr Pro Ile Ser Phe Ser Thr Asn Cys
                  100
                                      105
50
      Asp Ser Ile Ser Thr Ala Val
              115
      (2) INFORMATION FOR SEO ID NO: 428:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 102 amino acids
55
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
60
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..102
```

(D) OTHER INFORMATION: / Ceres Seq. ID 2032725

```
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:428:
      Met Ala Ser Pro Thr Glu Ser Thr Ile Gln Cys Gly Thr Val Thr Ser
                                           10
      Thr Leu Ala Gln Cys Leu Thr Tyr Leu Thr Asn Ser Gly Pro Leu Pro
 5
                                       25
      Ser Gln Cys Cys Val Gly Val Lys Ser Leu Tyr Gln Leu Ala Gln Thr
                                   40
      Thr Pro Asp Arg Lys Gln Val Cys Glu Cys Leu Lys Leu Ala Gly Lys
                                                   60
10
      Glu Ile Lys Gly Leu Asn Thr Asp Leu Val Ala Ala Leu Pro Thr Thr
                                               75
      Cys Gly Val Ser Ile Pro Tyr Pro Ile Ser Phe Ser Thr Asn Cys Asp
                                           .90
      Ser Ile Ser Thr Ala Val
15
                  100
      (2) INFORMATION FOR SEQ ID NO:429:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 75 amino acids
                (B) TYPE: amino acid
20
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MCLECULE TYPE: peptide
          (ix) FEATURE:
                 A) NAME/KEY: peptide
25
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 2032726
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:429:
      Lys Ala Ile Lys Asn Lys Asn Gly Val Tyr Ser Glu Asp His His Met
                                          10
30
      Pro His Cys Pro Tyr Asp Leu His Gly Ile Pro Asn Arg Val Asn His
                                      25
                  20
      Pro Val Trp Asp Ser Asp Glu His Thr Gly Thr Val Pro Asp Leu Leu
                                  40
      Asp Gln Gln Trp Ser Ile Ala Ile Thr Met Leu Arg Gly Ser Gln Val
35
                              55
      Ile Val Prc Ile Gly Ser Asp His Thr Gly Pro
                          7.0
      (2) INFORMATION FOR SEQ ID NO:430:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 198 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
45
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..198
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035536
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:430:
50
      atgtcgttgg tatggttgga agcgatgttg cctctcggaa tcatcggtgg gatgctctgt
                                                                             120
      atcatgggca attotcagta ctacatccac aaagcttatc atggccgtcc taagcacatc
      qqccacqatq aatqqqatqt tqctatqqaa aqacqcqaca aqaaaqtcqt cqaqaaaqct
      gcagctcctt cctcatga
      (2) INFORMATION FOR SEQ ID NO:431:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 65 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
60
          (ix) FEATURE:
                (A) NAME/KEY: peptide
```

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(B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035537
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:431:
      Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile Ile Gly
 5
                     5
                                          10
      Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His Lys Ala
                  20
                                      25
      Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp Val Ala
10
     Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Pro Ser
                              55
      Ser
      65
      (2) INFORMATION FOR SEQ ID NO:432:
15
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 65 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035538
25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:432:
      Cys Arg Trp Tyr Gly Trp Lys Arg Cys Cys Leu Ser Glu Ser Ser Val
                                         10
      Gly Cys Ser Val Ser Trp Ala Ile Leu Ser Thr Thr Ser Thr Lys Leu
                  20
                                      25
30
      Ile Met Ala Val Leu Ser Thr Ser Ala Thr Met Asn Gly Met Leu Leu
                                  40
      Trp Lys Asp Ala Thr Arg Lys Ser Ser Arg Lys Leu Gln Leu Leu Pro
                              55
      His
35
      65
      (2) INFORMATION FOR SEQ ID NO:433:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 57 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..57
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035539
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:433:
      Met Leu Pro Leu Gly Ile Ile Gly Gly Met Leu Cys Ile Met Gly Asn
                                          10
50
      Ser Gln Tyr Tyr Ile His Lys Ala Tyr His Gly Arg Pro Lys His Ile
                  20
                                      25
      Gly His Asp Glu Trp Asp Val Ala Met Glu Arg Arg Asp Lys Lys Val
                                  40
      Val Glu Lys Ala Ala Ala Pro Ser Ser
55
          50
                             5.5
      (2) INFORMATION FOR SEQ ID NO:434:
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 573 base pairs
                (B) TYPE: nucleic acid
60
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
```

550

```
(ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..573
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035575
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:434:
      atggagactt ctatgaggta taccagcaat tccaagtcta tgaagattca tgccaaagag
                                                                              120
      aaggttccgg tgaactcaaa aacccattta cagcttcatg gagagttaga tactggaact
      ggggctccga gttacttctg tgcqatqatt agacactttt ttcctqaggc ttcaacaggc
                                                                             180
      cttggggtag gattgcatta tgataagcgc caaaagcttc ggtgtcttgt acgcggaaaa
                                                                              240
10
      aaagagtttc ctgtaagagc tgataagcgt gtaaccttta atattaaagg gcggtgtgat
                                                                              300
      attgatcagg acttaaatca gaagaacccc aaaggagcag cagaatttgc ctggaacata
      atggatttca aggaagatca ggatgtacgg atcaaagttg gctacgaaat gtttgataag
                                                                             420
      gtcccttata tgcagattag agaaaacaat tggactctca acgcgaacat gaagggaaaa
                                                                              480
      tggaacttgc ggtatgacct gtaactgcat ttttttcaat catcatctga gaaatgtatt
                                                                             540
15
      gataccactg ctgatgaaca cattttaatt cta
      (2) INFORMATION FOR SEQ ID NO:435:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 167 amino acids
                (B) TYPE: amino acid
20
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
25
                (B) LOCATION: 1..167
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035576
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:435:
      Met Glu Thr Ser Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile
                      5
                                           10
                                                               15
30
      His Ala Lys Glu Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu
      His Gly Glu Leu Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala
                                  40
      Met Ile Arg His Phe Phe Pro Glu Ala Ser Thr Gly Leu Gly Val Gly
35
                              55
      Leu His Tyr Asp Lys Arg Gln Lys Leu Arg Cys Leu Val Arg Gly Lys
                                               75
      Lys Glu Phe Pro Val Arg Ala Asp Lys Arg Val Thr Phe Asn Ile Lys
40
      Gly Arg Cys Asp Ile Asp Gln Asp Leu Asn Gln Lys Asn Pro Lys Gly
                  100
                                       105
      Ala Ala Glu Phe Ala Trp Asn Ile Met Asp Phe Lys Glu Asp Gln Asp
                                  120
                                                       125
      Val Arg Ile Lys Val Gly Tyr Glu Met Phe Asp Lys Val Pro Tyr Met
45
          130
                              135
                                                   140
      Gln Ile Arg Glu Asn Asn Trp Thr Leu Asn Ala Asn Met Lys Gly Lys
                          150
                                               155
      Trp Asn Leu Arg Tyr Asp Leu
                      165
50
      (2) INFORMATION FOR SEQ ID NO:436:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 163 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
55
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..163
60
                (D) OTHER INFORMATION: / Ceres Seq. ID 2035577
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:436:
```

Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile His Ala Lys Glu

```
551
                                     10
     Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu His Gly Glu Leu
                        25
     Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala Met Ile Arg His
5
     Phe Phe Pro Glu Ala Ser Thr Gly Leu Gly Val Gly Leu His Tyr Asp
     Lys Arg Gln Lys Leu Arg Cys Leu Val Arg Gly Lys Lys Glu Phe Pro
                70
                                         75
     Val Arg Ala Asp Lys Arg Val Thr Phe Asn Ile Lys Gly Arg Cys Asp
10
                  85
                                     90
     Ile Asp Gln Asp Leu Asn Gln Lys Asn Pro Lys Gly Ala Ala Glu Phe
     Ala Trp Asn Ile Met Asp Phe Lys Glu Asp Gln Asp Val Arg Ile Lys
15
           115 120 125
     Val Gly Tyr Glu Met Phe Asp Lys Val Pro Tyr Met Gln Ile Arg Glu
      130 135
     Asn Asn Trp Thr Leu Asn Ala Asn Met Lys Gly Lys Trp Asn Leu Arg
                      150
                                         155
20
     Tyr Asp Leu
     (2) INFORMATION FOR SEQ ID NO:437:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 154 amino acids
25
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
30
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..154
              (D) OTHER INFORMATION: / Ceres Seq. ID 2035578
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:437:
     Met Lys Ile His Ala Lys Glu Lys Val Pro Val Asn Ser Lys Thr His
     1 5 10 15
Leu Gln Leu His Gly Glu Leu Asp Thr Gly Thr Gly Ala Pro Ser Tyr
35
                                 25
          20
     Phe Cys Ala Met Ile Arg His Phe Phe Pro Glu Ala Ser Thr Gly Leu
                              40
40
     Gly Val Gly Leu His Tyr Asp Lys Arg Gln Lys Leu Arg Cys Leu Val
                          55
     Arg Gly Lys Lys Glu Phe Pro Val Arg Ala Asp Lys Arg Val Thr Phe
                       70
     Asn Ile Lys Gly Arg Cys Asp Ile Asp Gln Asp Leu Asn Gln Lys Asn
45
                                      90
                  85
     Pro Lys Gly Ala Ala Glu Phe Ala Trp Asn Ile Met Asp Phe Lys Glu
              100 105
     Asp Gln Asp Val Arg Ile Lys Val Gly Tyr Glu Met Phe Asp Lys Val
            115 120 125
50
     Pro Tyr Met Gln Ile Arg Glu Asn Asn Trp Thr Leu Asn Ala Asn Met
        130 135 140
     Lys Gly Lys Trp Asn Leu Arg Tyr Asp Leu
             150
     (2) INFORMATION FOR SEQ ID NO:438:
55
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 750 base pairs
               (B) TYPE: nucleic acid
              (C) STRANDEDNESS: single
              (D) TOPOLOGY: linear
60
         (ii) MOLECULE TYPE: DNA (genomic)
         (ix) FEATURE:
              (A) NAME/KEY: -
```

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552
                (B) LOCATION: 1..750
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036457
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:438:
      atqacqactc agatcagcaa gaagagaaag tttgtagcgg acggtgtgtt ctacqctqaa
                                                                              60
 5
      ttgaatgagg ttcttacaag agagctagca gaggatggtt actctggtgt tqaqqttagq
                                                                             120
      qttactccta tqaqqactqa qattatcatc aqaqctacac qtactcaqaa tqttctcqqt
                                                                             180
      gagaagggga ggagaattag ggaattgact tecettgtte agaagagatt caagttteea
                                                                             240
      gttgacagtg ttgagcttta tgccgagaag gttaacaaca gaggtctctg tgccatcgct
                                                                             300
      caggetgagt etttaegtta caagettett ggtggteteg etgttegtag ggettgetat
                                                                             360
10
      qqtqtqttga qgtttgttat ggagagtgga gctaagggat gcgaggttat cgtgagtgga
                                                                             420
      aagettegtg etgeeagage caagtetatg aagtteaaag atggetaeat ggtgteatet
                                                                             480
      ggtcaaccaa ctaaggaata catagactct gcagtgagac atgttttgct tagacaaggt
                                                                             540
      gtgttgggaa tcaaggtgaa ggttatgctt gattgggacc ctaagggcat atcaggacca
                                                                             600
      aagacaccat tgcctgatgt tgtgatcatt cattctccta aagaagaaga ggccatctat
                                                                             660
15
      geacetgete aggitgetge eceggetget etegtageag atgeaceaet eacageegta
                                                                             720
      gattaccctg cgatgatccc agtcgcctaa
      (2) INFORMATION FOR SEQ ID NO:439:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 249 amino acids
20
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
25
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..249
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036458
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:439:
      Met Thr Thr Gln Ile Ser Lys Lys Arg Lys Phe Val Ala Asp Gly Val
30
                                          10
      Phe Tyr Ala Glu Leu Asn Glu Val Leu Thr Arg Glu Leu Ala Glu Asp
                  2.0
                                      25
      Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile
                                  4 0
35
      Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg
                              5.5
      Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys Arg Phe Lys Phe Pro
                          70
                                               7.5
      Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val Asn Asn Arg Gly Leu
40
                      85
                                          90
      Cvs Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly
                  100
                                      105
      Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu
                                  120
45
      Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala
                              135
                                                   140
      Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Val Ser Ser
                          150
                                              155
      Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ser Ala Val Arg His Val Leu
50
                                          170
                      165
      Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys Val Met Leu Asp Trp
                                      185
      Asp Pro Lys Gly Ile Ser Gly Pro Lys Thr Pro Leu Pro Asp Val Val
                                  200
                                                       205
55
      Ile Ile His Ser Pro Lys Glu Glu Glu Ala Ile Tyr Ala Pro Ala Gln
                              215
                                                   220
      Val Ala Ala Pro Ala Ala Leu Val Ala Asp Ala Pro Leu Thr Ala Val
                          230
                                               235
                                                                   240
      Asp Tyr Pro Ala Met Ile Pro Val Ala
60
                      245
      (2) INFORMATION FOR SEQ ID NO:440:
           (i) SEQUENCE CHARACTERISTICS:
```

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(A) LENGTH: 206 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..206
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036459
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:440:
     Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu
                                          10
      Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser Leu Val Gln Lys
                  20
                                      25
15
      Arg Phe Lys Phe Pro Val Asp Ser Val Glu Leu Tyr Ala Glu Lys Val
                                  40
      Asn Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr
      Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu
20
                          70
                                              75
      Arg Phe Val Met Glu Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser
                      85
                                          90
      Gly Lys Leu Arg Ala Ala Arg Ala Lys Ser Met Lys Phe Lys Asp Gly
                  100
                                      105
                                                          110
25
      Tyr Met Val Ser Ser Gly Gln Pro Thr Lys Glu Tyr Ile Asp Ser Ala
                                 120
             115
                                                      125
      Val Arg His Val Leu Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys
                             135
                                                  140
      Val Met Leu Asp Trp Asp Pro Lys Gly Ile Ser Gly Pro Lys Thr Pro
30
                         150
                                              155
      Leu Pro Asp Val Val Ile Ile His Ser Pro Lys Glu Glu Glu Ala Ile
                                          170
                      165
      Tyr Ala Pro Ala Gln Val Ala Ala Pro Ala Ala Leu Val Ala Asp Ala
                                      185
35
      Pro Leu Thr Ala Val Asp Tyr Pro Ala Met Ile Pro Val Ala
                                  200
                                                      205
      (2) INFORMATION FOR SEQ ID NO:441:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 549 base pairs
40
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
45
                (A) NAME/KEY: -
                (B) LOCATION: 1..549
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036585
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:441:
      atggttctca agacggagct ttgtcgtttc agtggacaga agatttaccc aggaagagga
                                                                              60
50
      attaggttta teegatetga tteteaggtt ttettgttte ttaacteaaa atgtaagagg
                                                                             120
      tacttccata acaagttgaa gccatccaag cttgcatgga ctgccatgta cagaaagcaa
                                                                             180
      cacaagaagg atgcagcaca agaggctgtg aagagaagga gacgtgccac caagaagcca
                                                                             240
      tactcaaggt ccattgttgg tgctaccttg gaagtaattc agaagaagag agctgagaag
                                                                             300
      cctgaagttc gtgatgcagc cagggaagct gctctgcgtg agatcaagga aagaatcaaa
                                                                             360
55
      aagaccaaag atgaaaagaa ggctaagaag gtggaatttg cttctaagca acagaaggtc
                                                                             420
      aaggetaatt teeceaaage tgetgetgea teeaagggte etaaggtggt ggtggeaaae
                                                                             480
      gctgaagage ttaaagccat cttttctcac tctgcgtctt ttctgctagt agctactttt
                                                                             540
      agtagttga
      (2) INFORMATION FOR SEQ ID NO:442:
60
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 182 amino acids
                (B) TYPE: amino acid
```

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(C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
 5
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..182
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036586
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:442:
     Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr
10
                                          10
      Pro Gly Arg Gly Ile Arg Phe Ile Arg Ser Asp Ser Gln Val Phe Leu
                  20
                                      25
      Phe Leu Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Lys Leu Lys Pro
                                  40
15
      Ser Lys Leu Ala Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp
                              55
      Ala Ala Gln Glu Ala Val Lys Arg Arg Arg Arg Ala Thr Lys Lys Pro
                          70
                                              75
      Tyr Ser Arg Ser Ile Val Gly Ala Thr Leu Glu Val Ile Gln Lys Lys
20
                      8.5
                                          90
      Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu
                  100
                                      105
                                                          110
      Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala
                                 120
             115
                                                      125
25
      Lys Lys Val Glu Phe Ala Ser Lys Gln Gln Lys Val Lys Ala Asn Phe
                             135
                                                 140
      Pro Lys Ala Ala Ala Ala Ser Lys Gly Pro Lys Val Val Val Ala Asn
                         150
                                             155
      Ala Glu Glu Leu Lys Ala Ile Phe Ser His Ser Ala Ser Phe Leu Leu
30
                     165
                                         170
      Val Ala Thr Phe Ser Ser
                  180
      (2) INFORMATION FOR SEQ ID NO: 443:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 127 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..127
                (D) OTHER INFORMATION: / Ceres Seq. ID 2036587
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:443:
45
      Met Tyr Arg Lys Gln His Lys Lys Asp Ala Ala Gln Glu Ala Val Lys
                                          10
      Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
                                      25
      Ala Thr Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val
50
              35
                                  40
      Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
                              55
      Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Val Glu Phe Ala Ser
                          70
                                               75
55
      Lys Gln Gln Lys Val Lys Ala Asn Phe Pro Lys Ala Ala Ala Ser
                                           90
                      85
      Lys Gly Pro Lys Val Val Ala Asn Ala Glu Glu Leu Lys Ala Ile
                                      105
                                                           110
      Phe Ser His Ser Ala Ser Phe Leu Leu Val Ala Thr Phe Ser Ser
60
                                  120
      (2) INFORMATION FOR SEQ ID NO:444:
           (i) SEQUENCE CHARACTERISTICS:
```

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555
                (A) LENGTH: 486 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..486
                (D) OTHER INFORMATION: / Ceres Seq. ID 2039554
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:444:
      atggctgtcg tcggcgctcc aatatcgtct ccggcggctc agctgcagac acaatttctc
                                                                              60
      tocaatocca ttotococcg otttogoogg totttotoca coggaaaato accagoaact
                                                                              120
      ttctccgtcg tagctatggc tccccagaaa aaggtgaaca aatatgatgc caagtggaag
                                                                              180
      aaacaatggt acggagctgg attgtttttc gaagggagtg agcaaataaa cgttgatgtt
                                                                              240
15
      ttcaagaagc tggagaagcg aaaagtgttg agcaacgttg agaaatctgg cctgctgtca
                                                                              300
      aaagcagagg ggttgggact cacattgtca tctcttgaga agcttaaagt cttctccaaa
                                                                              360
      gcagaggacc ttggtcttct cagtctcctt gagaacttag ctggaacatc gcctgcggtc
                                                                              420
      ttagcctcgg ctgcattacc agctctcacg agctgctatt gtagccgtgg tgttgatccc
                                                                              480
      ggatga
20
      (2) INFORMATION FOR SEQ ID NO:445:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 161 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..161
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 2039555
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:445:
      Met Ala Val Val Gly Ala Pro Ile Ser Ser Pro Ala Ala Gln Leu Gln
                                           10
      Thr Gln Phe Leu Ser Asn Pro Ile Leu Pro Arg Phe Arg Arg Ser Phe
35
                  20
                                       25
      Ser Thr Gly Lys Ser Pro Ala Thr Phe Ser Val Val Ala Met Ala Pro
                                   40
      Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys Trp Lys Lys Gln Trp Tyr
                              55
40
      Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu Gln Ile Asn Val Asp Val
                          70
                                               75
      Phe Lys Lys Leu Glu Lys Arg Lys Val Leu Ser Asn Val Glu Lys Ser
                                           90
                      8.5
      Gly Leu Leu Ser Lys Ala Glu Gly Leu Gly Leu Thr Leu Ser Ser Leu
45
                  100
                                       105
                                                           110
      Glu Lys Leu Lys Val Phe Ser Lys Ala Glu Asp Leu Gly Leu Leu Ser
                                  120
              115
                                                       125
      Leu Leu Glu Asn Leu Ala Gly Thr Ser Pro Ala Val Leu Ala Ser Ala
                              135
                                                   140
50
      Ala Leu Pro Ala Leu Thr Ser Cys Tyr Cys Ser Arg Gly Val Asp Pro
      145
                          150
                                               155
      Gly
      (2) INFORMATION FOR SEQ ID NO:446:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 116 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
```

(A) NAME/KEY: peptide

```
(B) LOCATION: 1..116
                (D) OTHER INFORMATION: / Ceres Seq. ID 2039556
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:446:
     Met Ala Pro Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys Trp Lys Lys
 5
                                        10
     Gln Trp Tyr Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu Gln Ile Asn
      Val Asp Val Phe Lys Lys Leu Glu Lys Arg Lys Val Leu Ser Asn Val
                                  40
10
      Glu Lys Ser Gly Leu Leu Ser Lys Ala Glu Gly Leu Gly Leu Thr Leu
      Ser Ser Leu Glu Lys Leu Lys Val Phe Ser Lys Ala Glu Asp Leu Glv
     Leu Leu Ser Leu Glu Asn Leu Ala Gly Thr Ser Pro Ala Val Leu
15
      Ala Ser Ala Ala Leu Pro Ala Leu Thr Ser Cys Tyr Cys Ser Arg Gly
                                      105
      Val Asp Pro Gly
              115
      (2) INFORMATION FOR SEQ ID NO:447:
20
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 378 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..378
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 2044283
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:447:
      atggcgaatt tgatgatgag attaccaatt agcttgagaa gcttctctgt ttcagcttct
                                                                              60
      tcatccaacg gttcgccgcc ggtgatcgga ggatctagcg gcggtgtagg accgatgatt
                                                                             120
      gtggaattac cgttggagaa gatacgaaga ccgttgatgc gaaccagatc caacgatcag
35
      aacaaagtga aagagcttat ggatagtatc cgtcaaatcg gtcttcaagt tccgattgat
                                                                             240
                                                                             300
      gtgattgaag ttgatggaac ttactatggg ttctcgggat gtcacagata cgaggcgcat
      cagaagctag ggcttccaac tatacgttgc aaaatccgta aaggaacaaa ggaaacatta
      aggcatcatc ttcgctga
      (2) INFORMATION FOR SEQ ID NO:448:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 125 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..125
                (D) OTHER INFORMATION: / Ceres Seq. ID 2044284
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:448:
      Met Ala Asn Leu Met Met Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser
                      5
                                          10
      Val Ser Ala Ser Ser Ser Asn Gly Ser Pro Pro Val Ile Gly Gly Ser
                                      25
55
      Ser Gly Gly Val Gly Pro Met Ile Val Glu Leu Pro Leu Glu Lys Ile
                                  40
      Arg Arg Pro Leu Met Arg Thr Arg Ser Asn Asp Gln Asn Lys Val Lys
                              55
      Glu Leu Met Asp Ser Ile Arg Gln Ile Gly Leu Gln Val Pro Ile Asp
60
                                               75
      Val Ile Glu Val Asp Gly Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg
```

WO 00/40695 557 Tyr Glu Ala His Gln Lys Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile 100 105 Arg Lys Gly Thr Lys Glu Thr Leu Arg His His Leu Arg 120 115 (2) INFORMATION FOR SEQ ID NO:449: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 121 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..121 (D) OTHER INFORMATION: / Ceres Seq. ID 2044285 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:449: Met Met Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser Val Ser Ala Ser 1.0 Ser Ser Asn Gly Ser Pro Pro Val Ile Gly Gly Ser Ser Gly Gly Val 20 25 Gly Pro Met Ile Val Glu Leu Pro Leu Glu Lys Ile Arg Arg Pro Leu 40 Met Arg Thr Arg Ser Asn Asp Gln Asn Lvs Val Lvs Glu Leu Met Asp 55 Ser Ile Arg Gln Ile Gly Leu Gln Val Pro Ile Asp Val Ile Glu Val 70 Asp Gly Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg Tyr Glu Ala His 85

Gln Lys Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile Arg Lys Gly Thr 100 105 Lys Glu Thr Leu Arg His His Leu Arg 115

(2) INFORMATION FOR SEQ ID NO:450:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 120 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- 40 (ix) FEATURE:

5

10

15

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25

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35

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- (A) NAME/KEY: peptide
- (B) LOCATION: 1..120
- (D) OTHER INFORMATION: / Ceres Seq. ID 2044286

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:450:

45 Met Arg Leu Pro Ile Ser Leu Arg Ser Phe Ser Val Ser Ala Ser Ser 10 Ser Asn Gly Ser Pro Pro Val Ile Gly Gly Ser Ser Gly Gly Val Gly 25

Pro Met Ile Val Glu Leu Pro Leu Glu Lys Ile Arg Arg Pro Leu Met 50 40

Arg Thr Arg Ser Asn Asp Gln Asn Lys Val Lys Glu Leu Met Asp Ser 55

Ile Arg Gln Ile Gly Leu Gln Val Pro Ile Asp Val Ile Glu Val Asp 75

55 Gly Thr Tyr Tyr Gly Phe Ser Gly Cys His Arg Tyr Glu Ala His Gln

Lys Leu Gly Leu Pro Thr Ile Arg Cys Lys Ile Arg Lys Gly Thr Lys 105

Glu Thr Leu Arg His His Leu Arg 115

(2) INFORMATION FOR SEQ ID NO:451:

(i) SEQUENCE CHARACTERISTICS:

```
558
                (A) LENGTH: 1236 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: ~
                (B) LOCATION: 1..1236
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048114
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:451:
      atgagaaqac ctagtcaaat gatgaggctt ctattaacat cctttttcqq tqttattqtt
                                                                              60
      qqtttcctta tqqqtattac ttttccaacc ttqactttaa ctaaqatqaa tcttccatcc
                                                                             120
      acattgtttc cctcgattga tcttgcatac attgaggata aatactctga catatcaaga
                                                                             180
      caaagactat ttggttcttg gtcttcgaca aaaggcctca aactcaagaa tgacatccct
                                                                             240
15
      gaccetecat ataactataa tgacactaag gttgatgata gaacgttega getattgeag
                                                                             300
      atatgggttt cgactaaccc ccgtggtgct gagaggctac caccagatat agtcacgcct
                                                                             360
      gaatcagatt tttacctccg tcgactgtgg ggcgacccta atgaggattt aacagtcaag
                                                                             420
      cagcggtatc tagtaacatt tacggttggc tatgatcaga ggaaaaatat agacactqtq
                                                                             480
      ttgaagaagt tctcagataa cttctctata atgctgtttc actacgatgg ccgggcaagc
                                                                             540
20
      qaatqqqaaq aqtttqaatq gtccaaqcqa qccattcatg tgaqcattcq qaaacaaaca
                                                                             600
      aaatggtggt acgctaagcg atttcttcat cctgacatag ttgcccccta tgaatatatc
                                                                             660
      ttcatatqqq atqaqqatct tqqcqtqqaa cactttqatt cqqaaaaata tctqqcqqtq
                                                                             720
      qtqaaqaaqc atggtttgga aatctcacag cctggattag agccatatga agggctcaca
                                                                             780
      tgggagatga ccaagaaaag agacgacact gaagtccaca agcatgctga ggaaaggaat
                                                                             840
25
      qggtggtgca ctgatcccaa tttaccccct tgtgcagcgt ttgtggagat tatggctcct
                                                                             900
      qttttctccc qcaaggcatg gcgctgtgtg tggcatatga ttcagaacga tttqattcat
                                                                             960
      qqatqqqqtc tqqactttqc cqttcqqaaa tqtqttcaqa acqcacacqa qaaaattqqa
                                                                            1020
      gttgtagatg ctcaatggat tatacatcaa ggtgttccat cattagggaa tcaaggacaa
                                                                            1080
      ccagagcaag ggaaacaacc atgggaaggg gtgagagaac gatgcaggag agagtggaca
                                                                            1140
30
      atgtttcaag acagattgga tgatgctgaa aaagcttatt ttgaagcatc tgctcacaag
                                                                            1200
      aatgcttctt cacggcctca cgggaattgg gtatag
      (2) INFORMATION FOR SEQ ID NO:452:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 411 amino acids
35
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..411
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048115
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:452:
      Met Arg Arg Pro Ser Gln Met Met Arg Leu Leu Thr Ser Phe Phe
45
                                           1.0
      Gly Val Ile Val Gly Phe Leu Met Gly Ile Thr Phe Pro Thr Leu Thr
                                       25
      Leu Thr Lys Met Asn Leu Pro Ser Thr Leu Phe Pro Ser Ile Asp Leu
                                   40
50
      Ala Tyr Ile Glu Asp Lys Tyr Ser Asp Ile Ser Arg Gln Arg Leu Phe
                              55
                                                   60
      Gly Ser Trp Ser Ser Thr Lys Gly Leu Lys Leu Lys Asn Asp Ile Pro
                          70
                                               75
      Asp Pro Pro Tyr Asn Tyr Asn Asp Thr Lys Val Asp Asp Arg Thr Phe
55
                                           90
      Glu Leu Leu Gln Ile Trp Val Ser Thr Asn Pro Arg Gly Ala Glu Arg
                                       105
      Leu Pro Pro Asp Ile Val Thr Pro Glu Ser Asp Phe Tyr Leu Arg Arg
                                   120
60
      Leu Trp Gly Asp Pro Asn Glu Asp Leu Thr Val Lys Gln Arg Tyr Leu
                               135
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Val Thr Phe Thr Val Gly Tyr Asp Gln Arg Lys Asn Ile Asp Thr Val

559 155 150 145 Leu Lys Lys Phe Ser Asp Asn Phe Ser Ile Met Leu Phe His Tyr Asp 165 170 175 Gly Arg Ala Ser Glu Trp Glu Glu Phe Glu Trp Ser Lys Arg Ala Ile 5 185 190 His Val Ser Ile Arg Lys Gln Thr Lys Trp Trp Tyr Ala Lys Arg Phe 200 Leu His Pro Asp Ile Val Ala Pro Tyr Glu Tyr Ile Phe Ile Trp Asp 215 10 Glu Asp Leu Gly Val Glu His Phe Asp Ser Glu Lys Tyr Leu Ala Val 230 235 240 Val Lys Lys His Gly Leu Glu Ile Ser Gln Pro Gly Leu Glu Pro Tyr 245 250 Glu Gly Leu Thr Trp Glu Met Thr Lys Lys Arg Asp Asp Thr Glu Val 15 260 265 His Lys His Ala Glu Glu Arg Asn Gly Trp Cys Thr Asp Pro Asn Leu 280 275 Pro Pro Cys Ala Ala Phe Val Glu Ile Met Ala Pro Val Phe Ser Arg 295 300 20 Lys Ala Trp Arg Cys Val Trp His Met Ile Gln Asn Asp Leu Ile His 310 315 Gly Trp Gly Leu Asp Phe Ala Val Arg Lys Cys Val Gln Asn Ala His 330 Glu Lys Ile Gly Val Val Asp Ala Gln Trp Ile Ile His Gln Gly Val 25 340 345 Pro Ser Leu Gly Asn Gln Gly Gln Pro Glu Gln Gly Lys Gln Pro Trp 360 Glu Gly Val Arg Glu Arg Cys Arg Arg Glu Trp Thr Met Phe Gln Asp 375 380 30 Arg Leu Asp Asp Ala Glu Lys Ala Tyr Phe Glu Ala Ser Ala His Lys 390 395 Asn Ala Ser Ser Arg Pro His Gly Asn Trp Val 405 410 (2) INFORMATION FOR SEQ ID NO:453: 35 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 405 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 40 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..405 (D) OTHER INFORMATION: / Ceres Seq. ID 2048116 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:453: Met Met Arg Leu Leu Thr Ser Phe Phe Gly Val Ile Val Gly Phe 10 Leu Met Gly Ile Thr Phe Pro Thr Leu Thr Leu Thr Lys Met Asn Leu 25 50 Pro Ser Thr Leu Phe Pro Ser Ile Asp Leu Ala Tyr Ile Glu Asp Lys 40 Tyr Ser Asp Ile Ser Arg Gln Arg Leu Phe Gly Ser Trp Ser Ser Thr Lys Gly Leu Lys Leu Lys Asn Asp Ile Pro Asp Pro Pro Tyr Asn Tyr 55 Asn Asp Thr Lys Val Asp Asp Arg Thr Phe Glu Leu Leu Gln Ile Trp 90 8.5 Val Ser Thr Asn Pro Arg Gly Ala Glu Arg Leu Pro Pro Asp Ile Val 105 Thr Pro Glu Ser Asp Phe Tyr Leu Arg Arg Leu Trp Gly Asp Pro Asn 60 120 Glu Asp Leu Thr Val Lys Gln Arg Tyr Leu Val Thr Phe Thr Val Gly PCT/US00/00466

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560
                            135
                                                 140
     Tyr Asp Gln Arg Lys Asn Ile Asp Thr Val Leu Lys Lys Phe Ser Asp
                    150
                                            155
     Asn Phe Ser Ile Met Leu Phe His Tyr Asp Gly Arg Ala Ser Glu Trp
 5
                                         170
                     165
     Glu Glu Phe Glu Trp Ser Lys Arg Ala Ile His Val Ser Ile Arg Lys
                 180
                                     185
     Gln Thr Lys Trp Trp Tyr Ala Lys Arg Phe Leu His Pro Asp Ile Val
                                200
                                                     205
10
     Ala Pro Tyr Glu Tyr Ile Phe Ile Trp Asp Glu Asp Leu Gly Val Glu
                             215
     His Phe Asp Ser Glu Lys Tyr Leu Ala Val Val Lys Lys His Gly Leu
                                             235
                         230
     Glu Ile Ser Gln Pro Gly Leu Glu Pro Tyr Glu Gly Leu Thr Trp Glu
15
                     245
                                         250
     Met Thr Lys Lys Arg Asp Asp Thr Glu Val His Lys His Ala Glu Glu
                                    265
                 260
     Arg Asn Gly Trp Cys Thr Asp Pro Asn Leu Pro Pro Cys Ala Ala Phe
             275 280
20
     Val Glu Ile Met Ala Pro Val Phe Ser Arg Lys Ala Trp Arg Cys Val
                            295
                                                 300
      Trp His Met Ile Gln Asn Asp Leu Ile His Gly Trp Gly Leu Asp Phe
                         310
                                             315
     Ala Val Arg Lys Cys Val Gln Asn Ala His Glu Lys Ile Gly Val Val
25
                     325
                                         330
     Asp Ala Gln Trp Ile Ile His Gln Gly Val Pro Ser Leu Gly Asn Gln
                 340
                                     345
      Gly Gln Pro Glu Gln Gly Lys Gln Pro Trp Glu Gly Val Arg Glu Arg
                                360
30
      Cys Arg Arg Glu Trp Thr Met Phe Gln Asp Arg Leu Asp Asp Ala Glu
                             375
                                                380
      Lys Ala Tyr Phe Glu Ala Ser Ala His Lys Asn Ala Ser Ser Arg Pro
                         390
                                             395
      His Gly Asn Trp Val
35
                     405
      (2) INFORMATION FOR SEQ ID NO:454:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 404 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..404
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048117
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:454:
      Met Arg Leu Leu Thr Ser Phe Phe Gly Val Ile Val Gly Phe Leu
                                         10
      Met Gly Ile Thr Phe Pro Thr Leu Thr Leu Thr Lys Met Asn Leu Pro
50
                                     25
                  20
      Ser Thr Leu Phe Pro Ser Ile Asp Leu Ala Tyr Ile Glu Asp Lys Tyr
                                  40
      Ser Asp Ile Ser Arg Gln Arg Leu Phe Gly Ser Trp Ser Ser Thr Lys
55
                              55
      Gly Leu Lys Leu Lys Asn Asp Ile Pro Asp Pro Pro Tyr Asn Tyr Asn
                                             75
      Asp Thr Lys Val Asp Asp Arg Thr Phe Glu Leu Leu Gln Ile Trp Val
                                          90
60
      Ser Thr Asn Pro Arg Gly Ala Glu Arg Leu Pro Pro Asp Ile Val Thr
                                     105
      Pro Glu Ser Asp Phe Tyr Leu Arg Arg Leu Trp Gly Asp Pro Asn Glu
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			115					120					125				
	Asp	Leu 130	Thr	Val	Lys	Gln	Arg 135	Tyr	Leu	Val	Thr	Phe 140	Thr	Val	Gly	Tyr	
5	Asp 145	Gln	Arg	Lys	Asn	Ile 150	Asp	Thr	Val	Leu	Lys 155	Lys	Phe	Ser	Asp	Asn 160	
	Phe	Ser	Ile	Met	Leu 165	Phe	His	Tyr	Asp	Gly 170	Arg	Ala	Ser	Glu	Trp 175	Glu	
	Glu	Phe	Glu	Trp 180	Ser	Lys	Arg	Ala	Ile 185	His	Val	Ser	Ile	Arg 190	Lys	Gln	
10	Thr	Lys	Trp 195	Trp	Tyr	Ala	Lys	Arg 200	Phe	Leu	His	Pro	Asp 205	Ile	Val	Ala	
	Pro	Tyr 210	Glu	Tyr	Ile	Phe	Ile 215	Trp	Asp	Glu	Asp	Leu 220	Gly	Val	Glu	His	
15	225	_				230		Ala			235					240	
0.0					245			Pro		250				_	255		
		_	_	260	_	_		Glu	265					270		-	
20			275					Asn 280					285				
		290					295	Ser				300	_	_		-	
25	305					310		Ile			315			_		320	
					325			Ala		330					335	_	
2.0				340				Gly	345					350		_	•
30			355		_	_		Pro 360					365		_	_	
		370		_			375	Gln				380	_			_	
35	385	-	Pne Trp		Ата	390	Ата	His	ьуs	ASN	395	ser	ser	Arg	Pro	ніs 400	
	(2)		-		FOR	SEO	TD I	NO:4	55:								
40	(2,		) SE	QUEN	CE C	HARA	CTER	ISTI ase p	CS:	S							
			(	B) T	YPE:	nuc	leic	acio	d								
		(ii	(	D) T	OPOL	OGY:	lin			c)							
45			) FE	ATUR A) N	E:			, ,									
			(	B) L	OCAT	:NOI	1	649 TION	: /	Cere	s Se	q. I	D 20	4827	1		
<b>5</b> 0			) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:45	5:					
50																ttccgg	60 120
															180		
	gaggtettag gttacttcaa teetttgeea ggeeaggaeg gtggtaagag gatgggtete 24														240		
55														300 360			
																cgtaaa gcagag	420
	aat	aaaa	cag	ttaa	tgcc	aa t	gata	acca	g cc	taag	gaag	agg	atac	aga	agcc	aagagt	480
	gcatgattca ttagccttct gtcatcgtag cttttcaagt tcactttgtt gtcgattata 540 ttgtgtaatg cagcattaga caactgactt gtttcctttg tttggcgata aacggcaagg 600																
60								gact caca							aacg	gcaagg	600
								NO:4		9		223	,				
		(i	) SE	QUEN	CE C	HARA	CTER	ISTI	CS:								

```
(A) LENGTH: 135 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..135
               (D) OTHER INFORMATION: / Ceres Seq. ID 2048272
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:456:
10
     Met Val Val Arg Ile Arg Leu Ser Arg Phe Gly Cys Lys Asn Arg Pro
                                        10
     Phe Phe Arg Val Met Ala Ala Asp Ser Arg Ser Pro Arg Asp Gly Lys
                                   25
15
     His Leu Glu Val Leu Gly Tyr Phe Asn Pro Leu Pro Gly Gln Asp Gly
                                40
     Gly Lys Arg Met Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser
                            55
     Val Gly Ala Gln Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser
20
                        70
                                            75
     Gly Leu Leu Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly
                    85 90
     Ala Arg Asp Thr Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp
                100 105
                                                       110
25
     Ala Glu Asn Lys Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu
            115 120
     Asp Thr Glu Ala Lys Ser Ala
         130
              135
      (2) INFORMATION FOR SEQ ID NO:457:
30
         (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 115 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
35
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..115
               (D) OTHER INFORMATION: / Ceres Seq. ID 2048273
40
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:457:
     Met Ala Ala Asp Ser Arg Ser Pro Arg Asp Gly Lys His Leu Glu Val
                                        10
      Leu Gly Tyr Phe Asn Pro Leu Pro Gly Gln Asp Gly Gly Lys Arg Met
                                    25
45
      Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser Val Gly Ala Gln
                                40
      Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser Gly Leu Leu Pro
                            55
      Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly Ala Arg Asp Thr
50
                                            75
                         7.0
      Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp Ala Glu Asn Lys
                                        90
                     85
      Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu Asp Thr Glu Ala
                 100
                                     105
55
      Lys Ser Ala
             115
      (2) INFORMATION FOR SEQ ID NO:458:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 84 amino acids
60
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
```

```
(ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..84
 5
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048274
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:458:
      Met Gly Leu Lys Phe Asp Arg Ile Lys Tyr Trp Leu Ser Val Gly Ala
                                           10
      Gln Pro Ser Asp Pro Val Gln Arg Leu Leu Phe Arg Ser Gly Leu Leu
10
                  20
                                       25
      Pro Pro Pro Met Val Ala Met Gly Arg Lys Gly Gly Ala Arg Asp
                                  40
      Thr Arg Pro Val Asp Pro Met Thr Gly Arg Tyr Val Asp Ala Glu Asn
                              55
                                                   60
15
      Lys Thr Val Asn Ala Asn Asp Asn Gln Pro Lys Glu Glu Asp Thr Glu
                          70
      Ala Lys Ser Ala
      (2) INFORMATION FOR SEQ ID NO: 459:
20
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 285 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
25
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..285
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048331
30
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:459:
      atgaaactgt gtaatcacca tactggaaag tttgtttcgg agagcgtcga agatcagact
                                                                              60
      gagcaggtac tcaaaaacat gggggagata ttgaaagcta gtggtgctga ttattcctcq
                                                                             120
      gtggtgaaga caacaatcat gttggctgat ttggctgact tcaagacagt gaacgagata
      tatgccaaat acttcccagc tectteteca geacgatega egtateaagt tgcagetttg
35
      cctctaaacq ccaagatcga gattgaatgt attgcaacac tctag
      (2) INFORMATION FOR SEQ ID NO: 460:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 94 amino acids
                (B) TYPE: amino acid
40
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
45
                (B) LOCATION: 1..94
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048332
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:460:
      Met Lys Leu Cys Asn His His Thr Gly Lys Phe Val Ser Glu Ser Val
                                           10
50
      Glu Asp Gln Thr Glu Gln Val Leu Lys Asn Met Gly Glu Ile Leu Lys
                  20
                                       25
      Ala Ser Gly Ala Asp Tyr Ser Ser Val Val Lys Thr Thr Ile Met Leu
                                   4.0
                                                       4.5
      Ala Asp Leu Ala Asp Phe Lys Thr Val Asn Glu Ile Tyr Ala Lys Tyr
55
                              55
                                                   60
      Phe Pro Ala Pro Ser Pro Ala Arg Ser Thr Tyr Gln Val Ala Ala Leu
                          70
                                               75
      Pro Leu Asn Ala Lys Ile Glu Ile Glu Cys Ile Ala Thr Leu
                                           90
                      85
60
      (2) INFORMATION FOR SEQ ID NO:461:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 68 amino acids
```

```
(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..68
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048333
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:461:
10
     Met Gly Glu Ile Leu Lys Ala Ser Gly Ala Asp Tyr Ser Ser Val Val
                                         10
      Lys Thr Thr Ile Met Leu Ala Asp Leu Ala Asp Phe Lys Thr Val Asn
                                      25
      Glu Ile Tyr Ala Lys Tyr Phe Pro Ala Pro Ser Pro Ala Arg Ser Thr
15
                              40
      Tyr Gln Val Ala Ala Leu Pro Leu Asn Ala Lys Ile Glu Ile Glu Cys
                              55
      Ile Ala Thr Leu
20
      (2) INFORMATION FOR SEQ ID NO: 462:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 48 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..48
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048334
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:462:
      Met Leu Ala Asp Leu Ala Asp Phe Lys Thr Val Asn Glu Ile Tyr Ala
                                          10
      Lys Tyr Phe Pro Ala Pro Ser Pro Ala Arg Ser Thr Tyr Gln Val Ala
35
               20
                                      25
      Ala Leu Pro Leu Asn Ala Lys Ile Glu Ile Glu Cys Ile Ala Thr Leu
                                 40
      (2) INFORMATION FOR SEQ ID NO:463:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 627 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
45
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..627
                (D) OTHER INFORMATION: / Ceres Seq. ID 2048466
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:463:
50
      atgatggaca agaatgaaaa ccctagtttt ttgctctctc gaagttccgt gcgactctct
                                                                              60
      gctcgagcaa gatctcctac cgatacaatg attatctcag aggctaaccg caaagaaatc
                                                                             120
      tgcaaqtacc tcttcaaaga aggagtttgc tttgcgaaga aggatttcaa tcttgctaag
                                                                             180
      catccqttga tcgatgtccc caacctacaa gtgattaagc ttatgcagag tttcaaatcc
                                                                             240
      aaqqaqtatq ttaqqqaqac atttqcctqq atqcattact attqqttttt qaccaatqaa
                                                                             300
55
      qqqatcqaqt tettqaqaac ttatettaat ettecatetg atgttqttee tqetaetttq
                                                                             360
      aagaagtcag ctaagcctgg tggtcgtcca tttggtggcc cacctggtga tcgctcaaga
                                                                             420
      ggacctcgcc atgaaggagg agaccgtccc aggttttggtg accgtgatgg gtaccgtgca
                                                                             480
      ggtcctcgag ctggtggtga gtttggaggt gaaaagggtg gagctcccgc agattaccag
                                                                             540
      ccatctttcc aaggaagtgg ccgtggtttt ggccgtggtg ctggtggcta cagcgcagct
                                                                             600
60
      gcaccatctg gttcaggttt gccttga
      (2) INFORMATION FOR SEQ ID NO: 464:
```

(i) SEQUENCE CHARACTERISTICS:

```
(A) LENGTH: 208 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
 5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..208
              (D) OTHER INFORMATION: / Ceres Seq. ID 2048467
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:464:
     Met Met Asp Lys Asn Glu Asn Pro Ser Phe Leu Leu Ser Arg Ser Ser
                                      10
     Val Arg Leu Ser Ala Arg Ala Arg Ser Pro Thr Asp Thr Met Ile Ile
                20
                                   25
15
     Ser Glu Ala Asn Arg Lys Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly
                               40
     Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Ala Lys His Pro Leu Ile
                           55
     Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser
20
                                          75
     Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe
                                   90
                   8.5
     Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro
               100 105
25
     Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly
           115 120
     Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Ser Arg Gly Pro Arg His
                                             140
        130 135
     Glu Gly Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Ala
30
                                          155
                      150
     Gly Pro Arg Ala Gly Gly Glu Phe Gly Gly Glu Lys Gly Gly Ala Pro
                                     170
                    165
     Ala Asp Tyr Gln Pro Ser Phe Gln Gly Ser Gly Arg Gly Phe Gly Arg
                180 185
35
     Gly Ala Gly Gly Tyr Ser Ala Ala Ala Pro Ser Gly Ser Gly Leu Pro
                     200
      (2) INFORMATION FOR SEQ ID NO: 465:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 207 amino acids
40
               (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
45
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..207
              (D) OTHER INFORMATION: / Ceres Seq. ID 2048468
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:465:
     Met Asp Lys Asn Glu Asn Pro Ser Phe Leu Leu Ser Arg Ser Ser Val
50
                                       10
     Arg Leu Ser Ala Arg Ala Arg Ser Pro Thr Asp Thr Met Ile Ile Ser
                                   25
     Glu Ala Asn Arg Lys Glu Ile Cys Lys Tyr Leu Phe Lys Glu Gly Val
                               40
55
     Cys Phe Ala Lys Lys Asp Phe Asn Leu Ala Lys His Pro Leu Ile Asp
                            55
     Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser Lys
                                          75
                        7.0
     Glu Tyr Val Arq Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe Leu
60
                                       90
     Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro Ser
```

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566
     Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly Arg
                                 120
     Pro Phe Gly Gly Pro Pro Gly Asp Arg Ser Arg Gly Pro Arg His Glu
                                               140
                             135
 5
     Gly Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Ala Gly
                        150
                                            155
     Pro Arg Ala Gly Gly Glu Phe Gly Gly Glu Lys Gly Gly Ala Pro Ala
                                        170
                    165
     Asp Tyr Gln Pro Ser Phe Gln Gly Ser Gly Arg Gly Phe Gly Arg Gly
10
                180
                         185
     Ala Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser Gly Leu Pro
                      200
      (2) INFORMATION FOR SEQ ID NO: 466:
          (i) SEQUENCE CHARACTERISTICS:
15
               (A) LENGTH: 179 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
20
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..179
               (D) OTHER INFORMATION: / Ceres Seq. ID 2048469
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:466:
25
     Met Ile Ile Ser Glu Ala Asn Arg Lys Glu Ile Cys Lys Tyr Leu Phe
                                        10
     Lys Glu Gly Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Ala Lys His
                 20
                                    25
      Pro Leu Ile Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser
30
                                40
      Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr
                             5.5
     Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu
                                            75
35
     Asn Leu Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys
                                         90
     Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Ser Arg Gly
                                    105
     Pro Arg His Glu Gly Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly
40
            115
                                120
     Tyr Arg Ala Gly Pro Arg Ala Gly Gly Glu Phe Gly Gly Glu Lys Gly
                            135
     Gly Ala Pro Ala Asp Tyr Gln Pro Ser Phe Gln Gly Ser Gly Arg Gly
                        150 155
45
     Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser
                                        170
     Gly Leu Pro
      (2) INFORMATION FOR SEQ ID NO: 467:
50
         (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 585 base pairs
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
55
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
               (A) NAME/KEY: -
               (B) LOCATION: 1..585
               (D) OTHER INFORMATION: / Ceres Seq. ID 2050485
60
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:467:
     atcttcggaa agtctcattt ctcgatcccc aattcgtgga ttagggttaa aagaaccatt
```

tttattctcq tcqcqcaaca acaaatccaq atcqaaaaaq gaagaagaga tcqaaatqqc

567

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gttgagaagg gtttacagtg aaatcagagg gaagaaggtg acggagcttc caggctatat
                                                                             180
      caaatcgact ttttcaatgg agaccgtgaa gacctctgtg aagagaggac tcgataacta
                                                                              240
      caacgaaaaa tacattcaga ccagctccgt tgatcctatc cttcatatct gcttctacgg
                                                                             300
      catggettte tettacettg tegeteteee taatgagegt egecatettg ageateagea
                                                                             360
      gcatgctaag gagcacggtg gtcattgatc tcgtgggaat cgtttcgatc tcgagatgat
                                                                             420
     tttagggggg ttgctgtgaa atctttctct gcttgatggt gacgacgact caagaattgt
                                                                             480
      gtcttattgt ttcgttttct tgaattttcc tggataatgt tgacctaaag gaaaaccttt
                                                                             540
      ctttcgaatt acactccatg atagtcaata attgaagcat catga
      (2) INFORMATION FOR SEQ ID NO: 468:
10
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 128 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
15
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..128
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050486
20
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:468:
      Ser Ser Glu Ser Leu Ile Ser Arg Ser Pro Ile Arg Gly Leu Gly Leu
                                           10
      Lys Glu Pro Phe Leu Phe Ser Ser Arg Asn Asn Lys Ser Arg Ser Lys
                  20
                                      2.5
                                                           30
25
      Lys Glu Glu Glu Ile Glu Met Ala Leu Arg Arg Val Tyr Ser Glu Ile
                                  40
                                                       45
      Arg Gly Lys Lys Val Thr Glu Leu Pro Gly Tyr Ile Lys Ser Thr Phe
                              55
                                                   60
      Ser Met Glu Thr Val Lys Thr Ser Val Lys Arg Gly Leu Asp Asn Tyr
30
                          70
                                               75
      Asn Glu Lys Tyr Ile Gln Thr Ser Ser Val Asp Pro Ile Leu His Ile
                                           90
      Cys Phe Tyr Gly Met Ala Phe Ser Tyr Leu Val Ala Leu Pro Asn Glu
                                      105
35
      Arg Arg His Leu Glu His Gln Gln His Ala Lys Glu His Gly Gly His
              115
                                  120
      (2) INFORMATION FOR SEQ ID NO: 469:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 90 amino acids
40
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
45
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..90
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050487
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:469:
      Met Ala Leu Arg Arg Val Tyr Ser Glu Ile Arg Gly Lys Lys Val Thr
50
                                           10
      Glu Leu Pro Gly Tyr Ile Lys Ser Thr Phe Ser Met Glu Thr Val Lys
                                       25
      Thr Ser Val Lys Arg Gly Leu Asp Asn Tyr Asn Glu Lys Tyr Ile Gln
                                   40
55
      Thr Ser Ser Val Asp Pro Ile Leu His Ile Cys Phe Tyr Gly Met Ala
      Phe Ser Tyr Leu Val Ala Leu Pro Asn Glu Arg Arg His Leu Glu His
                          70
                                               75
      Gln Gln His Ala Lys Glu His Gly Gly His
60
                      85
      (2) INFORMATION FOR SEQ ID NO: 470:
```

(i) SEOUENCE CHARACTERISTICS:

```
(A) LENGTH: 516 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..516
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050708
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:470:
      caaccaaaac aaaacataaa aaacaagtgg aagctttaaa acgagaggga gagagcaaaa
                                                                              60
      atggcgacgt cgggaacgta cgtgacggaa gttccqctaa aaggatcggc cgagaaatac
                                                                              120
      tacaagaggt ggaagaacga gaaccatgte tteeetgatg etateggeea ecacatecaa
                                                                              180
      aatgttaccg ttcacgaagg cgaacatgac tctcacgggt ctatcaggag ttggaactac
                                                                              240
15
                                                                              300
      acatgggatg gaaaggagga ggtgttcaag gagagaagag agatagacga tgagaccaaa
                                                                              360
      acgttgacgt taagaggact tgagggtcac gtgatggagc agctcaaagt gtacgacgtc
                                                                              420
      gtctaccaat tcattcccaa atctgaggat acctgcatcg gcaaaatcac tttaatatgg
      gagaagegea acgatgatte eccagaacea agegggtaca tgaaattegt caagagettg
                                                                              480
      qttqctqaca tqqqaaacca cqttaqcaaa acttaa
20
      (2) INFORMATION FOR SEQ ID NO:471:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 151 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..151
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050709
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:471:
      Met Ala Thr Ser Gly Thr Tyr Val Thr Glu Val Pro Leu Lys Gly Ser
                      5
                                           10
                                                               15
      Ala Glu Lys Tyr Tyr Lys Arg Trp Lys Asn Glu Asn His Val Phe Pro
35
                  20
                                       25
      Asp Ala Ile Gly His His Ile Gln Asn Val Thr Val His Glu Gly Glu
                                   40
      His Asp Ser His Gly Ser Ile Arg Ser Trp Asn Tyr Thr Trp Asp Gly
                               55
40
      Lys Glu Glu Val Phe Lys Glu Arg Glu Ile Asp Asp Glu Thr Lys
                          70
                                                                    8.0
      65
                                               75
      Thr Leu Thr Leu Arg Gly Leu Glu Gly His Val Met Glu Gln Leu Lys
                      85
                                           90
      Val Tyr Asp Val Val Tyr Gln Phe Ile Pro Lys Ser Glu Asp Thr Cys
45
                  100
                                       105
      Ile Gly Lys Ile Thr Leu Ile Trp Glu Lys Arg Asn Asp Asp Ser Pro
              115
                                   120
      Glu Pro Ser Gly Tyr Met Lys Phe Val Lys Ser Leu Val Ala Asp Met
                               135
50
      Gly Asn His Val Ser Lys Thr
      145
                           150
      (2) INFORMATION FOR SEQ ID NO:472:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1730 base pairs
55
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
60
                 (A) NAME/KEY: -
                 (B) LOCATION: 1..1730
                 (D) OTHER INFORMATION: / Ceres Seq. ID 2050901
```

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:472:
      atcgcgtgtt actcctttct aaaggaacca ttgtatatca tggaaggctt gacttacttg
                                                                              60
      aagcattett geteteeaag ggatteaegg tteettetea getgaattet ettgagtaeg
                                                                             120
      ctatggagat acttcagaac atccgtgatc cctacgaaaa cgccaacatt gctcttccag
                                                                             180
 5
      accactgece tgaaagtaaa aaacagaace aaaaacagag cattgttega tataaaaget
                                                                             240
      cgagaatcac cgaaataagc cttctctcta gcaggttttg gaagatcata taccgtacaa
                                                                             300
      qgcaqttqct tctaacaaac atcttagaat ctcttqtaqt cqqtcttqtc ttaqqcacta
                                                                             360
      tctaccttaa tatcggaact ggcaaagaag gaatcaggaa acgatttggc cttttcgcat
                                                                             420
      tcaccctcac attcctcctc tcctccacta cccaaaccct tccaatattc attgatgaac
                                                                             480
10
      gacctattct tctccgagaa acctcaagcg gactctacag actctcctct cacattcttg
                                                                             540
      caaacacttt ggttttcttg ccatacttgc tactcatcgc aatcatctac tctgtctcac
                                                                             600
      tctatttcct tgtaggactc tgcttttcat ggcaagctct cgcctacttt gtgctcgtaa
                                                                             660
      totggatcat tgtoctaatg gotaactott ttgtactttt ottgagotot otogoacota
                                                                             720
                                                                             780
      actacattgc tggaacatct tcagtgacca ttcttcttgc ggctttcttc ttgttctctg
15
      gttacttcat ctctaaagag agtcttccca agtactggct cttcatgtac ttcttctcaa
                                                                             840
      tgtacaagta tgcgttggac gcacttctga taaatgagta ctcgtgtctg cacaacaagt
                                                                             900
                                                                             960
      qcctggtctg gtttgaggaa gcttctgtga atagctgctt agttactgga ggtgacgtgt
      taqacaaqaa tqqqcttcat gaqaqacaqa qqatatatqt qttqqqacqt ccqtcattqq
                                                                            1020
      tatccggttt caaacttgag aaacaaggaa tacgtttctt gagaagcaag aaaaccctat
                                                                            1080
20
      tgctcccctt tgctcaagct cagtgcctct ttgcagcgaa agctcaacca acaagacgat
                                                                            1140
      ggaattatoo gagtattaat tgtgatottg ttogtoottt tatoagatta totoagagaa
                                                                            1200
      caatcqcaqa qaqatctqca aatacctttt caaaqctaqt qqaaccaqaq qqaqtatqct
      ttgctaagaa ggatttcaat ctcccaaagc atccgttgat tgatgtacca aacctgcaag
      tgattaaget catgeagagt tteaagteea aggagtaegt tagggagaea tttgegtgga
25
      tgcattatta ttggtttctg actaatgaag gaattgagtt cttgagaact tatcttaacc
      ttoottooga tgttgtooct gotactttga agaagtotgo taagoooggt ggtogtooct
                                                                            1500
      ttqqtqqccc acctqqtqat cqccaaaqaq qaccacctcq ctctqatqqa qaccqtccca
                                                                            1560
      gatttggtga ccgtgatgga taccgtggag gcccacgtgg tggtgatgag aagggtggag
                                                                            1620
      ctccagctga tttccagccg tctttccaag gaggtggtgg taggcctggt tttggccgtg
                                                                            1680
30
      qtqcaqqcqq ttacaqtqca qcaqcaccat ctqqttcaqq qttcccttqa
      (2) INFORMATION FOR SEQ ID NO:473:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 575 amino acids
                (B) TYPE: amino acid
35
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
40
                (B) LOCATION: 1..575
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050902
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:473:
      Arg Val Leu Leu Ser Lys Gly Thr Ile Val Tyr His Gly Arg Leu
                      5
                                           10
45
      Asp Leu Leu Glu Ala Phe Leu Leu Ser Lys Gly Phe Thr Val Pro Ser
                  20
                                      25
      Gln Leu Asn Ser Leu Glu Tyr Ala Met Glu Ile Leu Gln Asn Ile Arg
                                  40
      Asp Pro Tyr Glu Asn Ala Asn Ile Ala Leu Pro Asp His Cys Pro Glu
50
                              55
      Ser Lys Lys Gln Asn Gln Lys Gln Ser Ile Val Arg Tyr Lys Ser Ser
                          70
      Arg Ile Thr Glu Ile Ser Leu Leu Ser Ser Arg Phe Trp Lys Ile Ile
55
      Tyr Arg Thr Arg Gln Leu Leu Thr Asn Ile Leu Glu Ser Leu Val
                  100
                                      105
                                                           110
      Val Gly Leu Val Leu Gly Thr Ile Tyr Leu Asn Ile Gly Thr Gly Lys
              115
                                  120
                                                       125
      Glu Gly Ile Arg Lys Arg Phe Gly Leu Phe Ala Phe Thr Leu Thr Phe
60
                              135
                                                   140
      Leu Leu Ser Ser Thr Thr Gln Thr Leu Pro Ile Phe Ile Asp Glu Arg
      145
                          150
                                               155
                                                                   160
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570
    Pro Ile Leu Leu Arg Glu Thr Ser Ser Gly Leu Tyr Arg Leu Ser Ser
    $165$ $170$ $175$ His Ile Leu Ala Asn Thr Leu Val Phe Leu Pro Tyr Leu Leu Ile
                  185
 5
    Ala Ile Ile Tyr Ser Val Ser Leu Tyr Phe Leu Val Gly Leu Cys Phe
          195
               200
    Ser Trp Gln Ala Leu Ala Tyr Phe Val Leu Val Ile Trp Ile Ile Val
            215 220
    Leu Met Ala Asn Ser Phe Val Leu Phe Leu Ser Ser Leu Ala Pro Asn
10
         230 235 240
    Tyr Ile Ala Gly Thr Ser Ser Val Thr Ile Leu Leu Ala Ala Phe Phe
                 245 250
    Leu Phe Ser Gly Tyr Phe Ile Ser Lys Glu Ser Leu Pro Lys Tyr Trp
                              265 270
              260
15
    Leu Phe Met Tyr Phe Phe Ser Met Tyr Lys Tyr Ala Leu Asp Ala Leu
                                           285
                           280
    Leu Ile Asn Glu Tyr Ser Cys Leu His Asn Lys Cys Leu Val Trp Phe
                        295 300
    Glu Glu Ala Ser Val Asn Ser Cys Leu Val Thr Gly Gly Asp Val Leu
20
        310
                                     315
    Asp Lys Asn Gly Leu His Glu Arg Gln Arg Ile Tyr Val Leu Gly Arg
                 325
                                 330
     Fro Ser Leu Val Ser Gly Phe Lys Leu Glu Lys Gln Gly Ile Arg Phe
                              345
25
    Leu Arg Ser Lys Lys Thr Leu Leu Pro Phe Ala Gln Ala Gln Cys
                           360
     Leu Phe Ala Ala Lys Ala Gln Pro Thr Arg Arg Trp Asn Tyr Pro Ser
                        375
                                        380
     Ile Asn Cys Asp Leu Val Arg Pro Phe Ile Arg Leu Ser Gln Arg Thr
30
                     390
                                     395
    Ile Ala Glu Arg Ser Ala Asn Thr Phe Ser Lys Leu Val Glu Pro Glu
                 405 410
    Gly Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Pro Lys His Pro Leu
                              425 430
              420
     Ile Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser Phe Lys
35
                           440
     Ser Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr Tyr Trp
                        455
                                        460
     Phe Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu
40
                     470
                                     475
     Pro Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly
                                  490 495
                 485
     Gly Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Gln Arg Gly Pro Pro
                               505
45
    Arg Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg
                           520
     Gly Gly Pro Arg Gly Gly Asp Glu Lys Gly Gly Ala Pro Ala Asp Phe
                        535
     Gln Pro Ser Phe Gln Gly Gly Gly Arg Pro Gly Phe Gly Arg Gly
50
     545 550
                                     555 560
    Ala Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser Gly Phe Pro
                565
                                 570
     (2) INFORMATION FOR SEQ ID NO:474:
         (i) SEQUENCE CHARACTERISTICS:
55
             (A) LENGTH: 535 amino acids
             (B) TYPE: amino acid
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
60
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..535
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571
              (D) OTHER INFORMATION: / Ceres Seq. ID 2050903
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:474:
     Met Glu Ile Leu Gln Asn Ile Arg Asp Pro Tyr Glu Asn Ala Asn Ile
                                      10
 5
     Ala Leu Pro Asp His Cys Pro Glu Ser Lys Lys Gln Asn Gln Lys Gln
                20
                                  25
     Ser Ile Val Arg Tyr Lys Ser Ser Arg Ile Thr Glu Ile Ser Leu Leu
                               4.0
     Ser Ser Arg Phe Trp Lys Ile Ile Tyr Arg Thr Arg Gln Leu Leu
10
                           55
                                              60
     Thr Asn Ile Leu Glu Ser Leu Val Val Gly Leu Val Leu Gly Thr Ile
                        70
     Tyr Leu Asn Ile Gly Thr Gly Lys Glu Gly Ile Arg Lys Arg Phe Gly
                    85
                                       90
15
     Leu Phe Ala Phe Thr Leu Thr Phe Leu Leu Ser Ser Thr Thr Gln Thr
                100
                                  105
     Leu Pro Ile Phe Ile Asp Glu Arg Pro Ile Leu Leu Arg Glu Thr Ser
                               120
     Ser Gly Leu Tyr Arg Leu Ser Ser His Ile Leu Ala Asn Thr Leu Val
20
                           135
                                              140
     Phe Leu Pro Tyr Leu Leu Ile Ala Ile Ile Tyr Ser Val Ser Leu
                        150
                                          155
     Tyr Phe Leu Val Gly Leu Cys Phe Ser Trp Gln Ala Leu Ala Tyr Phe
                                      170 175
                    165
25
     Val Leu Val Ile Trp Ile Ile Val Leu Met Ala Asn Ser Phe Val Leu
                180
                                   185
     Phe Leu Ser Ser Leu Ala Pro Asn Tyr Ile Ala Gly Thr Ser Ser Val
             195
                               200
     Thr Ile Leu Leu Ala Ala Phe Phe Leu Phe Ser Gly Tyr Phe Ile Ser
30
                           215
                                              220
     Lys Glu Ser Leu Pro Lys Tyr Trp Leu Phe Met Tyr Phe Phe Ser Met
                        230
                                          235
     Tyr Lys Tyr Ala Leu Asp Ala Leu Leu Ile Asn Glu Tyr Ser Cys Leu
                    245
                                       250
35
     His Asn Lys Cys Leu Val Trp Phe Glu Glu Ala Ser Val Asn Ser Cys
                260
                                   265
     Leu Val Thr Gly Gly Asp Val Leu Asp Lys Asn Gly Leu His Glu Arg
                               280
     Gln Arg Ile Tyr Val Leu Gly Arg Pro Ser Leu Val Ser Gly Phe Lys
40
                           295
     Leu Glu Lys Gln Gly Ile Arg Phe Leu Arg Ser Lys Lys Thr Leu Leu
                        310
                                          315
     Leu Pro Phe Ala Gln Ala Gln Cys Leu Phe Ala Ala Lys Ala Gln Pro
                    325
                                       330
45
     Thr Arg Arg Trp Asn Tyr Pro Ser Ile Asn Cys Asp Leu Val Arg Pro
                                  345
     Phe Ile Arg Leu Ser Gln Arg Thr Ile Ala Glu Arg Ser Ala Asn Thr
                               360
     Phe Ser Lys Leu Val Glu Pro Glu Gly Val Cys Phe Ala Lys Lys Asp
50
                           375
                                              380
     Phe Asn Leu Pro Lys His Pro Leu Ile Asp Val Pro Asn Leu Gln Val
                        390
                                         395
     Ile Lys Leu Met Gln Ser Phe Lys Ser Lys Glu Tyr Val Arg Glu Thr
                    405
                                      410
55
     Phe Ala Trp Met His Tyr Tyr Trp Phe Leu Thr Asn Glu Gly Ile Glu
                420
                                  425
     Phe Leu Arg Thr Tyr Leu Asn Leu Pro Ser Asp Val Pro Ala Thr
                              440
     Leu Lys Lys Ser Ala Lys Pro Gly Gly Arg Pro Phe Gly Gly Pro Pro
60
       450 455
                                 460
     Gly Asp Arg Gln Arg Gly Pro Pro Arg Ser Asp Gly Asp Arg Pro Arg
     465
                        470
                                          475
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572 Phe Gly Asp Arg Asp Gly Tyr Arg Gly Gly Pro Arg Gly Gly Asp Glu 490 Lys Gly Gly Ala Pro Ala Asp Phe Gln Pro Ser Phe Gln Gly Gly 505 510 Gly Arg Pro Gly Phe Gly Arg Gly Ala Gly Gly Tyr Ser Ala Ala Ala 515 520 525 Pro Ser Gly Ser Gly Phe Pro 530 (2) INFORMATION FOR SEQ ID NO:475: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 350 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..350 (D) OTHER INFORMATION: / Ceres Seq. ID 2050904 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:475: Met Ala Asn Ser Phe Val Leu Phe Leu Ser Ser Leu Ala Pro Asn Tyr 1 5 10 Ile Ala Gly Thr Ser Ser Val Thr Ile Leu Leu Ala Ala Phe Phe Leu 20 25 25 Phe Ser Gly Tyr Phe Ile Ser Lys Glu Ser Leu Pro Lys Tyr Trp Leu 40 Phe Met Tyr Phe Phe Ser Met Tyr Lys Tyr Ala Leu Asp Ala Leu Leu 55 Ile Asn Glu Tyr Ser Cys Leu His Asn Lys Cys Leu Val Trp Phe Glu 75 30 70 Glu Ala Ser Val Asn Ser Cys Leu Val Thr Gly Gly Asp Val Leu Asp 90 Lys Asn Gly Leu His Glu Arg Gln Arg Ile Tyr V l Leu Gly Arg Pro 105 35 Ser Leu Val Ser Gly Phe Lys Leu Glu Lys Gln Gly Ile Arg Phe Leu 120 Arg Ser Lys Lys Thr Leu Leu Pro Phe Ala Gln Ala Gln Cys Leu 135 Phe Ala Ala Lys Ala Gln Pro Thr Arg Arg Trp Asn Tyr Pro Ser Ile 40 150 155 Asn Cys Asp Leu Val Arg Pro Phe Ile Arg Leu Ser Gln Arg Thr Ile 165 170 175 Ala Glu Arg Ser Ala Asn Thr Phe Ser Lys Leu Val Glu Pro Glu Gly 180 185 190 Val Cys Phe Ala Lys Lys Asp Phe Asn Leu Pro Lys His Pro Leu Ile 45 200 205 Asp Val Pro Asn Leu Gln Val Ile Lys Leu Met Gln Ser Phe Lys Ser 210 215 220 Lys Glu Tyr Val Arg Glu Thr Phe Ala Trp Met His Tyr Tyr Trp Phe 50 230 235 Leu Thr Asn Glu Gly Ile Glu Phe Leu Arg Thr Tyr Leu Asn Leu Pro 245 250 255 Ser Asp Val Val Pro Ala Thr Leu Lys Lys Ser Ala Lys Pro Gly Gly 260 265 270 55 Arg Pro Phe Gly Gly Pro Pro Gly Asp Arg Gln Arg Gly Pro Pro Arg 275 280 285 Ser Asp Gly Asp Arg Pro Arg Phe Gly Asp Arg Asp Gly Tyr Arg Gly 295 300 Gly Pro Arg Gly Gly Asp Glu Lys Gly Gly Ala Pro Ala Asp Phe Gln 60 305 310 315 Pro Ser Phe Gln Gly Gly Gly Arg Pro Gly Phe Gly Arg Gly Ala 325 330

573 Gly Gly Tyr Ser Ala Ala Pro Ser Gly Ser Gly Phe Pro 340 345 (2) INFORMATION FOR SEQ ID NO:476: (i) SEQUENCE CHARACTERISTICS: 5 (A) LENGTH: 535 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 10 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..535 (D) OTHER INFORMATION: / Ceres Seq. ID 2051325 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:476: 15 gctacataac tcaaagtaac aaacatacac atataacata tagaaagatc gaaaacaaaa gatcgggatg agtatggctc caaagacctc aactacactt gctttattcc ttgtgaccaa tattetette etcaacetea ttaccetgag ttgegeagae aataettgee caagagaegt totcaaactt togacatgot ogaatgttot caacctcato aacttgaago toggggcaco 240 agetatgagg cettgttget etattetett tggtetaatt gatetegatg ttgetgtttg 20 cettigeace gegeteaage teageettet iggeateace alegaeacte etaiteacet taacttggct cttaacgcct gtggaggtac ccttcctgat ggattccgtt gcccaacata 420 gctacaatat attcatattt tctatctatg ttcttgaatt ggttttgtct aacatccaaa tttcaaacct aaattcatgt ataagcaaat aacaagtaaa attaagactt ttatg (2) INFORMATION FOR SEQ ID NO:477: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 117 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..117 (D) OTHER INFORMATION: / Ceres Seq. ID 2051326 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:477: Met Ser Met Ala Pro Lys Thr Ser Thr Thr Leu Ala Leu Phe Leu Val 5 10 Thr Asn Ile Leu Phe Leu Asn Leu Ile Thr Leu Ser Cys Ala Asp Asn 25 40 Thr Cys Pro Arg Asp Val Leu Lys Leu Ser Thr Cys Ser Asn Val Leu 40 Asn Leu Ile Asn Leu Lys Leu Gly Ala Pro Ala Met Arg Pro Cys Cys 55 Ser Ile Leu Phe Gly Leu Ile Asp Leu Asp Val Ala Val Cys Leu Cys 45 75 70 Thr Ala Leu Lys Leu Ser Leu Leu Gly Ile Thr Ile Asp Thr Pro Ile 85 90 His Leu Asn Leu Ala Leu Asn Ala Cys Gly Gly Thr Leu Pro Asp Gly 100 105 50 Phe Arg Cys Pro Thr 115 (2) INFORMATION FOR SEQ ID NO:478: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 115 amino acids (B) TYPE: amino acid 55 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 60 (A) NAME/KEY: peptide (B) LOCATION: 1..115

(D) OTHER INFORMATION: / Ceres Seq. ID 2051327

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:478:
     Met Ala Pro Lys Thr Ser Thr Thr Leu Ala Leu Phe Leu Val Thr Asn
                                          10
     Ile Leu Phe Leu Asn Leu Ile Thr Leu Ser Cys Ala Asp Asn Thr Cys
 5
                                      25
     Pro Arg Asp Val Leu Lys Leu Ser Thr Cys Ser Asn Val Leu Asn Leu
                                  40
                                                      45
     Ile Asn Leu Lys Leu Gly Ala Pro Ala Met Arg Pro Cys Cys Ser Ile
                              55
10
     Leu Phe Gly Leu Ile Asp Leu Asp Val Ala Val Cys Leu Cys Thr Ala
                          70
                                              75
     Leu Lys Leu Ser Leu Leu Gly Ile Thr Ile Asp Thr Pro Ile His Leu
                     8.5
                                          90
     Asn Leu Ala Leu Asn Ala Cys Gly Gly Thr Leu Pro Asp Gly Phe Arg
1.5
                                                          110
                 100
                                      105
     Cys Pro Thr
              115
      (2) INFORMATION FOR SEQ ID NO:479:
           (i) SEQUENCE CHARACTERISTICS:
20
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          ii) MOLECULE TYPE: peptide
25
          lix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 2051328
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:479:
30
     Met Leu Leu Phe Ala Phe Ala Pro Arg Ser Ser Ser Ala Phe Leu Ala
                  5
                                          10
      Ser Pro Ser Thr Leu Leu Phe Thr Leu Thr Trp Leu Leu Thr Pro Val
                  20
                                      25
      Glu Val Pro Phe Leu Met Asp Ser Val Ala Gln His Ser Tyr Asn Ile
35
                                  40
      Phe Ile Phe Ser Ile Tvr Val Leu Glu Leu Val Leu Ser Asn Ile Gln
                              55
      Ile Ser Asn Leu Asn Ser Cys Ile Ser Lys
                          70
40
      (2) INFORMATION FOR SEQ ID NO:480:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 517 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..517
50
                (D) OTHER INFORMATION: / Ceres Seq. ID 2051633
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:480:
      cataaccctc ttqqaaaqaq tctcaacact tqcaqaqaaa aaqaacaaqq aaqatcccqq
                                                                             60
      aaaatggcaa cggcgattgt acgttcagct ctttcccgag cagtgactcg cgcagctccg
                                                                            120
      aagacatccg tcgctcctaa gcgaaacttt tcctcttccg ccggccatga cgatgcttat
                                                                            180
55
      gaagetgega agtgggagaa gataaettat etgggtattg etagttgeae tgetetaget
                                                                            240
      gtctatgttt tatccaaggg ccatcatcac ggcgaagacc ctcctgccta tccgcatatg
                                                                            300
      cacatccgca acaaggagtt tccttggggt ccggatggtc tgtttgaggt gaagcacaac
                                                                            360
      aaagagcact gagtcttgcg tggtcataat aacgtcttct tggtttattt gaaaggctaa
                                                                            420
      aatgttttac cgtatttgtt ctcaccgttt gtcaacgatt tgctactcca atctcttttc
                                                                            480
60
      ttttgttggg aaataaaagt taatactttg cttggtc
      (2) INFORMATION FOR SEQ ID NO:481:
           (i) SEQUENCE CHARACTERISTICS:
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575 (A) LENGTH: 123 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..123 (D) OTHER INFORMATION: / Ceres Seq. ID 2051634 10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:481: His Asn Pro Leu Gly Lys Ser Leu Asn Thr Cys Arg Glu Lys Glu Gln 10 Gly Arg Ser Arg Lys Met Ala Thr Ala Ile Val Arg Ser Ala Leu Ser 20 25 15 Arg Ala Val Thr Arg Ala Ala Pro Lys Thr Ser Val Ala Pro Lys Arg 40 Asn Phe Ser Ser Ala Gly His Asp Asp Ala Tyr Glu Ala Ala Lys 55 60 Trp Glu Lys Ile Thr Tyr Leu Gly Ile Ala Ser Cys Thr Ala Leu Ala 20 70 75 Val Tyr Val Leu Ser Lys Gly His His His Gly Glu Asp Pro Pro Ala 90 Tyr Pro His Met His Ile Arg Asn Lys Glu Phe Pro Trp Gly Pro Asp 105 100 25 Gly Leu Phe Glu Val Lys His Asn Lys Glu His 115 120 (2) INFORMATION FOR SEQ ID NO:482: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 102 amino acids 30 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 35 (A) NAME/KEY: peptide (B) LOCATION: 1..102 (D) OTHER INFORMATION: / Ceres Seq. ID 2051635 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:482: Met Ala Thr Ala Ile Val Arg Ser Ala Leu Ser Arg Ala Val Thr Arg 40 10 Ala Ala Pro Lys Thr Ser Val Ala Pro Lys Arg Asn Phe Ser Ser Ser 20 25 Ala Gly His Asp Asp Ala Tyr Glu Ala Ala Lys Trp Glu Lys Ile Thr 40 45 Tyr Leu Gly Ile Ala Ser Cys Thr Ala Leu Ala Val Tyr Val Leu Ser Lys Gly His His His Gly Glu Asp Pro Pro Ala Tyr Pro His Met His 70 75 Ile Arg Asn Lys Glu Phe Pro Trp Gly Pro Asp Gly Leu Phe Glu Val 50 85 90 Lys His Asn Lys Glu His 100 (2) INFORMATION FOR SEQ ID NO:483: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 410 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)

(ix) FEATURE:

(A) NAME/KEY: -(B) LOCATION: 1..410

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(D) OTHER INFORMATION: / Ceres Seq. ID 2051906
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:483:
      ttccttcttt ttcacatcta cccaaaattc caaaacacca tcgatttttt qctctcttga
                                                                               60
      aatgtgtctt tttatgtcca attcttctct tccaactaaa ccgaaccgga aaaccaggtt
                                                                              120
 5
     cggagatcgg tgtttactga tggccaaaca gcaacgaacc cgactttaca tacttcgacg
                                                                             180
      atgtgtctcc atgctgcttt gctggcacga ccactctatt tctgattagt ttgattcatg
                                                                              240
      aaccggacgg accaagaagt agaaatcaga agaatgtgat gtcatttaga gtttttcttg
                                                                              300
      ttttctttta gaactcttca aggaatcttt tgtaattgaa gaagactcta aggattgggc
                                                                             360
      cttttgggcc tttgtacata ttgtgtaaat aacatatgaa cttttttac
10
      (2) INFORMATION FOR SEQ ID NO:484:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 75 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
15
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..75
20
                (D) OTHER INFORMATION: / Ceres Seq. ID 2051907
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:484:
      Ser Phe Phe Phe Thr Ser Thr Gln Asn Ser Lys Thr Pro Ser Ile Phe
                      5
                                           10
      Cys Ser Leu Glu Met Cys Leu Phe Met Ser Asn Ser Ser Leu Pro Thr
25
                  20
                                                           30
                                       25
      Lys Pro Asn Arg Lys Thr Arg Phe Gly Asp Arg Cys Leu Leu Met Ala
                                   40
      Lys Gln Gln Arg Thr Arg Leu Tyr Ile Leu Arg Arg Cys Val Ser Met
                              55
30
      Leu Leu Cys Trp His Asp His Ser Ile Ser Asp
                          70
      (2) INFORMATION FOR SEQ ID NO:485:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 55 amino acids
35
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..55
                (D) OTHER INFORMATION: / Ceres Seq. ID 2051908
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:485:
      Met Cys Leu Phe Met Ser Asn Ser Ser Leu Pro Thr Lys Pro Asn Arg
45
                      5
                                           10
      Lys Thr Arg Phe Gly Asp Arg Cys Leu Leu Met Ala Lys Gln Gln Arg
                                       25
                                                           30
      Thr Arg Leu Tyr Ile Leu Arg Arg Cys Val Ser Met Leu Leu Cys Trp
              35
                                   40
50
      His Asp His Ser Ile Ser Asp
                               55
      (2) INFORMATION FOR SEQ ID NO:486:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 51 amino acids
55
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
60
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..51
                 (D) OTHER INFORMATION: / Ceres Seq. ID 2051909
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577 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:486: Met Ser Asn Ser Ser Leu Pro Thr Lys Pro Asn Arg Lys Thr Arg Phe 10 Gly Asp Arg Cys Leu Leu Met Ala Lys Gln Gln Arg Thr Arg Leu Tyr 5 20 25 Ile Leu Arg Arg Cys Val Ser Met Leu Leu Cys Trp His Asp His Ser 40 Ile Ser Asp 50 10 (2) INFORMATION FOR SEQ ID NO:487: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 719 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..719 2.0 (D) OTHER INFORMATION: / Ceres Seq. ID 2052403 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:487: acaaatcatt caaaqacata caaaataatt qaqttttttt tttaattaga aacaaatqqq gttgagtggt gttcttcatg tggaggttga ggrtaagtct ccggctgaaa agttctgggt 120 agcccttggc gacggcatca atctcttccc caaagctttc cctaacgact acaaaaccat 180 25 ccaaqttcta qccqqcqatq qcaacqctcc tqqctccatt cqcctcatta cttatqqaqa 240 aggateteca etggtgaaga tateggegga gaggategaa geagtggatt tggagaacaa 300 aagcatgtcg tacagcatca ttggcggaga aatgttggag tactacaaaa ccttcaaagg 360 aaccatcacc gttatcccta aggatggtgg tagccttctg aaatggtctg gtgagtttga 420 gaagaccgcc catgagatcg atgatccaca tgtcatcaag gactttgctg tcaagaactt 480 30 caaaqagata gatgagtato ttottaagca aactagtgoo taacactaga acctttaaat 540 tatataagag ggttcgaccg tctctataag atttttctaa ttaagaagtt gaataaagtg 600 gaaccccttt atgaatatcc acgttttatg atttcggagt ttatgcagcc tagtaggcca 660 taagcttttt acaaaagcca atttagtcga acattttgaa aaaatcgaac cttgtggta (2) INFORMATION FOR SEQ ID NO:488: 35 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 155 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 40 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..155 (D) OTHER INFORMATION: / Ceres Seq. ID 2052404 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:488: Met Gly Leu Ser Gly Val Leu His Val Glu Val Lys Ser Pro 10 Ala Glu Lys Phe Trp Val Ala Leu Gly Asp Gly Ile Asn Leu Phe Pro 25 50 Lys Ala Phe Pro Asn Asp Tyr Lys Thr Ile Gln Val Leu Ala Gly Asp 40 Gly Asn Ala Pro Gly Ser Ile Arg Leu Ile Thr Tyr Gly Glu Gly Ser 55 Pro Leu Val Lys Ile Ser Ala Glu Arq Ile Glu Ala Val Asp Leu Glu 55 70 75 Asn Lys Ser Met Ser Tyr Ser Ile Ile Gly Gly Glu Met Leu Glu Tyr Tyr Lys Thr Phe Lys Gly Thr Ile Thr Val Ile Pro Lys Asp Gly Gly 105

Ser Leu Leu Lys Trp Ser Gly Glu Phe Glu Lys Thr Ala His Glu Ile 115 120 125 Asp Asp Pro His Val Ile Lys Asp Phe Ala Val Lys Asn Phe Lys Glu

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578
         130
                            135
     Ile Asp Glu Tyr Leu Leu Lys Gln Thr Ser Ala
              150
      (2) INFORMATION FOR SEQ ID NO:489:
 5
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 739 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
10
          (iii) MOLECULE TYPE: DNA (genomic)
          'ix) FEATURE:
               (A) NAME/KEY: -
                (B) LOCATION: 1..739
                (D) OTHER INFORMATION: / Ceres Seq. ID 2053545
15
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:489:
     attittagtae attgttgace atcttttteg tatagactae tatetetgat etettgegag
     ttaagtcagt aactaggaaa attcagaagc qctctcaatc tcaaaaatat ccatggcggc
      gattacagaa tttctaccaa aagagtacgg atatgtcgtt ctcgtcctcg tcttctactg
      tttcctcaac ctctggatgg gtgctcaagt cggcagaget cgcaaaaggt acaacgtccc
                                                                           240
20
      ptatccaact statatgcaa tagaatsaga aaasaaagat gstaagstot tsaastgtgt
                                                                           300
      teagalagga cateaaaaet etttagagat gatgeeaatg tattteatae tgatgateet
                                                                           360
      aggaggaatg aagcaccott gratotgtac tggcottggt tagotttaca acgttagcog
                                                                           420
      attettetae tttaaaqqtt atqetactqq aqateecatq aaqeqtetta eqateqqqaa
                                                                           480
      atacqqtttc ttqqqqttqc taqqtctqat qatatqtacc atctcqtttq qtqtcactct
                                                                           540
25
     gatcottgct tgagctactc gtttctgggg ttaatgattc tctggtttgc tcgaagaata
                                                                           600
      tagaaccaat gottgtaago tgtocacaaa acttgtgtaa tactttagag tttgtcactt
                                                                           660
      ttaaaaagttt gtaataaatc atggcttcat agaacagttg aaatttcaca tccgtagacg
                                                                           720
      ttaataaaga tttgaatta
      (2) INFORMATION FOR SEQ ID NO:490:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 146 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..146
                (D) OTHER INFORMATION: / Ceres Seq. ID 2053546
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          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:490:
      Met Ala Ala Ile Thr Glu Phe Leu Pro Lys Glu Tyr Gly Tyr Val Val
                                         10
      Leu Val Leu Val Phe Tyr Cys Phe Leu Asn Leu Trp Met Gly Ala Gln
                                      25
                  20
45
      Val Gly Arg Ala Arg Lys Arg Tyr Asn Val Pro Tyr Pro Thr Leu Tyr
                                 40
                                                      45
      Ala Ile Glu Ser Glu Asn Lys Asp Ala Lys Leu Phe Asn Cys Val Gln
                             55
                                                  60
      Arg Gly His Gln Asn Ser Leu Glu Met Met Pro Met Tyr Phe Ile Leu
50
                         7.0
                                              75
      Met Ile Leu Gly Gly Met Lys His Pro Cys Ile Cys Thr Gly Leu Gly
                      85
                                         90
      Leu Leu Tyr Asn Val Ser Arg Phe Phe Tyr Phe Lys Gly Tyr Ala Thr
                                      105
                                                          110
55
      Gly Asp Pro Met Lys Arg Leu Thr Ile Gly Lys Tyr Gly Phe Leu Gly
                     120
                                          125
      Leu Leu Gly Leu Met Ile Cys Thr Ile Ser Phe Gly Val Thr Leu Ile
         130
                             135
      Leu Ala
60
      (2) INFORMATION FOR SEQ ID NO:491:
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(i) SEQUENCE CHARACTERISTICS:

WO 00/40695 579 (A) LENGTH: 118 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..118 (D) OTHER INFORMATION: / Ceres Seq. ID 2053547 10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:491: Met Gly Ala Gln Val Gly Arg Ala Arg Lys Arg Tyr Asn Val Pro Tyr 5 10 Pro Thr Leu Tyr Ala Ile Glu Ser Glu Asn Lys Asp Ala Lys Leu Phe 25 15 Asn Cys Val Gln Arg Gly His Gln Asn Ser Leu Glu Met Met Pro Met 40 Tyr Phe Ile Leu Met Ile Leu Gly Gly Met Lys His Pro Cys Ile Cys 55 Thr Gly Leu Gly Leu Leu Tyr Asn Val Ser Arg Phe Phe Tyr Phe Lys 20 75 Gly Tyr Ala Thr Gly Asp Pro Met Lys Arg Leu Thr Ile Gly Lys Tyr 85 90 Gly Phe Leu Gly Leu Gly Leu Met Ile Cys Thr Ile Ser Phe Gly 100 105 25 Val Thr Leu Ile Leu Ala 115 (2) INFORMATION FOR SEQ ID NO:492: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 74 amino acids (B) TYPE: amino acid 30 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 35 (A) NAME/KEY: peptide (B) LOCATION: 1..74 (D) OTHER INFORMATION: / Ceres Seq. ID 2053548 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:492: Met Met Pro Met Tyr Phe Ile Leu Met Ile Leu Gly Gly Met Lys His 40 10 Pro Cys Ile Cys Thr Gly Leu Gly Leu Leu Tyr Asn Val Ser Arg Phe 25 Phe Tyr Phe Lys Gly Tyr Ala Thr Gly Asp Pro Met Lys Arg Leu Thr 40 45 45 Ile Gly Lys Tyr Gly Phe Leu Gly Leu Leu Gly Leu Met Ile Cys Thr 55 Ile Ser Phe Gly Val Thr Leu Ile Leu Ala 70 (2) INFORMATION FOR SEQ ID NO:493: 50 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 681 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 55 (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:493: ccatccaaaa aaaacaaaac aaaaaattat attcaagaga aaaaggaaaa aatgaatttc

(ix) FEATURE:

60

(A) NAME/KEY: -(B) LOCATION: 1..681

atctccgatc aggtaaagaa actctcaagc tcaacaccag aggagccaga ccacaacaag

60 120

(D) OTHER INFORMATION: / Ceres Seq. ID 2053884

PCT/US00/00466 WO 00/40695 580 ccagtcgaag gaaccgaaac agctacaaga ccagctacca acgccgagct catggcaagt 180 gccaaggttg tagctgaagc tgctcaagcc gcagctcgta acgaatcaga caaactcgac 240 aagggtaaag tcgccggagc ctctgctgat atcttagacg ctgccgagaa atacggtaag 300 ttcgatgaaa agagtagcac tggtcagtac ctcgacaagg ctgagaagta tctcaacgac 360 tacgagtcgt cacactccac cggtgctggt ggtcctcctc ctccgacgag tcaggctgag 420 ccagcaagtc agcctgagcc ggcggctaaq aaagacgatg aagagtctgg tggtqggctt 480 ggaggttatg ccaagatggc tcaaggtttc ttgaagtgat ttgatcttta attgttgttc 540 atcattttcg taataataaa ttaaataact agtatcgttt gtgactagtt tatgttgctt 600 cgtttatgtt tatggggagt gacgagtgag tgtaataact tctggtgatc atgaatctaa 660 10 tccatctttg ttgtgattat g (2) INFORMATION FOR SEQ ID NO:494: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 172 amino acids (B) TYPE: amino acid 15 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 20 (B) LOCATION: 1..172 (D) OTHER INFORMATION: / Ceres Seq. ID 2053885 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:494: Pro Ser Lys Lys Asn Lys Thr Lys Asn Tyr Ile Gln Glu Lys Lys Glu 10 15 25 Lys Met Asn Phe Ile Ser Asp Gln Val Lys Lvs Leu Ser Ser Thr 20 25 Pro Glu Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala 40 Thr Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val 30 55 Ala Glu Ala Ala Gln Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp 75 Lys Gly Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ala Glu 90 35 Lys Tyr Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly Gln Tyr Leu Asp 105 110 Lys Ala Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly 120 125 Ala Gly Gly Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln 40 135 140 Pro Glu Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Leu 150 155 Gly Gly Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys 165 45 (2) INFORMATION FOR SEQ ID NO:495: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 155 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..155 55 (D) OTHER INFORMATION: / Ceres Seq. ID 2053886 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:495: Met Asn Phe Ile Ser Asp Gln Val Lys Lys Leu Ser Ser Ser Thr Pro 10

Glu Glu Pro Asp His Asn Lys Pro Val Glu Gly Thr Glu Thr Ala Thr

25 Arg Pro Ala Thr Asn Ala Glu Leu Met Ala Ser Ala Lys Val Val Ala 40

45

581 Glu Ala Ala Gln Ala Ala Arg Asn Glu Ser Asp Lys Leu Asp Lys 55 Gly Lys Val Ala Gly Ala Ser Ala Asp Ile Leu Asp Ala Ala Glu Lys 70 75 5 Tyr Gly Lys Phe Asp Glu Lys Ser Ser Thr Gly Gln Tyr Leu Asp Lys 90 Ala Glu Lys Tyr Leu Asn Asp Tyr Glu Ser Ser His Ser Thr Gly Ala 105 Gly Gly Pro Pro Pro Pro Thr Ser Gln Ala Glu Pro Ala Ser Gln Pro 10 120 Glu Pro Ala Ala Lys Lys Asp Asp Glu Glu Ser Gly Gly Gly Leu Gly 135 140 Gly Tyr Ala Lys Met Ala Gln Gly Phe Leu Lys 150 15 (2) INFORMATION FOR SEQ ID NO:496: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 115 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..115 25 (D) OTHER INFORMATION: / Ceres Seq. ID 2053887 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:496: Met Ala Ser Ala Lys Val Val Ala Glu Ala Ala Gln Ala Ala Ala Arg 1.0 Asn Glu Ser Asp Lys Leu Asp Lys Gly Lys Val Ala Gly Ala Ser Ala 30 25 Asp Ile Leu Asp Ala Ala Glu Lys Tyr Gly Lys Phe Asp Glu Lys Ser 40 Ser Thr Gly Gln Tyr Leu Asp Lys Ala Glu Lys Tyr Leu Asn Asp Tyr 5.5 60 35 Glu Ser Ser His Ser Thr Gly Ala Gly Gly Pro Pro Pro Pro Thr Ser 75 7.0 Gln Ala Glu Pro Ala Ser Gln Pro Glu Pro Ala Ala Lys Lys Asp Asp 90 Glu Glu Ser Gly Gly Leu Gly Gly Tyr Ala Lys Met Ala Gln Gly 40 100 105 Phe Leu Lys 115 (2) INFORMATION FOR SEO ID NO:497: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 797 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 50 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..797 (D) OTHER INFORMATION: / Ceres Seq. ID 2053908 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:497: 55 ttctctcaaa atccataaaa agagagaga ataaataaag agaaaactaa agaagctaga agatggagaa aagtaatgat catgacaagg ctagccacgg cggctccggt ggtggtgcca 120 cggagaagtg ggaggagact agcctcggaa tccgaactgc cgagacaatg ctccggttag ctccggtggg cctttgtgtt gcagcacttg ttgtcatgct taaggactct gagactaatg 240 agttcggttc aatttcttac tccaatctca cagcctttag gtacttggtg cacgcaaatg 300 60 gaatatgtgc aggctactct cttctatcag cagccattgc agcgatgcct cgttcttctt 360 cgacaatgcc tcgtqtttgg accttctttt qtctcqacca gcttctgacc tacctgqttc 420 ttgctgctgg agctgtatca gctgaggttc tatacttggc ttacaatgga gactcagcca 480

582 ttacttggag cgatgcatgt agctcctatg gcggtttctg tcatagagcc actgcttctg ttataatcac attetttqtq qtttqtttct acateqttct etetetaate teetettata 600 agetetttae tegetttgat eeteetteea ttgttgaete egeeaagaat etegaagteg ctgtcttcgg aagttagatc ctccatttgc tcttcaagta catctagttt gtcatgtttc 5 aaatqttqtg tgtttttaac tttggtcaag agaaagaatg cttatgtgtt tgctcttctt tcattgcttt tcttcct (2) INFORMATION FOR SEQ ID NO:498: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 204 amino acids 10 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide :ix) FEATURE: 15 (A) NAME/KEY: peptide (B) LOCATION: 1..204 (D) OTHER INFORMATION: / Ceres Seq. ID 2053909 xi) SEQUENCE DESCRIPTION: SEQ ID NO:498: Met Glu Lys Ser Asn Asp His Asp Lys Ala Ser His Gly Gly Ser Gly 20 10 Gly Gly Ala Thr Glu Lys Trp Glu Glu Thr Ser Leu Gly Ile Arg Thr 20 25 Ala Glu Thr Met Leu Arg Leu Ala Pro Val Gly Leu Cys Val Ala Ala 40 45 25 Leu Val Val Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile 55 Ser Tyr Ser Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly 70 75 Ile Cys Ala Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro 30 85 90 Arg Ser Ser Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp 105 100 110 Gln Leu Leu Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu 120 125 35 Val Leu Tyr Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp 135 140 Ala Cys Ser Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val 150 155 Ile Ile Thr Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile 40 165 170 Ser Ser Tyr Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp 180 185 Ser Ala Lys Asn Leu Glu Val Ala Val Phe Gly Ser 45 (2) INFORMATION FOR SEQ ID NO:499: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 169 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..169 55 (D) OTHER INFORMATION: / Ceres Seq. ID 2053910 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:499: Met Leu Arg Leu Ala Pro Val Gly Leu Cys Val Ala Ala Leu Val Val 10 Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile Ser Tyr Ser 25 60

Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly Ile Cys Ala

40

583 Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro Arg Ser Ser 55 Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp Gln Leu Leu 75 5 Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu Val Leu Tyr 90 Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp Ala Cys Ser 105 Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val Ile Ile Thr 10 115 120 Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile Ser Ser Tyr 135 Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp Ser Ala Lys 150 155 15 Asn Leu Glu Val Ala Val Phe Gly Ser 165 (2) INFORMATION FOR SEQ ID NO:500: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 153 amino acids 20 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 25 (A) NAME/KEY: peptide (B) LOCATION: 1..153 (D) OTHER INFORMATION: / Ceres Seq. ID 2053911 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:500: Met Leu Lys Asp Ser Glu Thr Asn Glu Phe Gly Ser Ile Ser Tyr Ser 30 10 Asn Leu Thr Ala Phe Arg Tyr Leu Val His Ala Asn Gly Ile Cys Ala 25 Gly Tyr Ser Leu Leu Ser Ala Ala Ile Ala Ala Met Pro Arg Ser Ser 40 35 Ser Thr Met Pro Arg Val Trp Thr Phe Phe Cys Leu Asp Gln Leu Leu 55 Thr Tyr Leu Val Leu Ala Ala Gly Ala Val Ser Ala Glu Val Leu Tyr 70 Leu Ala Tyr Asn Gly Asp Ser Ala Ile Thr Trp Ser Asp Ala Cys Ser 40 Ser Tyr Gly Gly Phe Cys His Arg Ala Thr Ala Ser Val Ile Ile Thr 105 Phe Phe Val Val Cys Phe Tyr Ile Val Leu Ser Leu Ile Ser Ser Tyr 120 45 Lys Leu Phe Thr Arg Phe Asp Pro Pro Ser Ile Val Asp Ser Ala Lys 135 Asn Leu Glu Val Ala Val Phe Gly Ser 150 (2) INFORMATION FOR SEQ ID NO:501: 50 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 652 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 55 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..652 (D) OTHER INFORMATION: / Ceres Seq. ID 2056123 60 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:501: attttttctt tctctctct ctccctaagc aaaactaaaa caagctatgg ctggtatgct tocoggagtt gagtgtgcaa ggaggcggcg ottocacggt ggtgctooto cgattgaato

60

584

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ctcgaacaca gcttctgtgg cgqctgcggc ggqacacqtc tggacacggc gaccatcgtt
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     ctctctttac actaccaatc atgagagcca ccaggcccat gtctccttct cggagagaag
                                                                            240
     tgttaggaat aaatcttatg gagaagacaa cgatgagaaa cttgacggag cagccaaaga
                                                                            300
     ggcaaagcag aggcttaaca agcggctgag aatcccacca cgtacaaggc aaaatggtaa
                                                                            360
 5
     agacaaaggg aataaattgg agcaaggaaa gggtaaacct ctcgggggact taccgaccga
                                                                            420
     ggtggtcggg ttaaagaaga gccgaggaag gttgatggaa tggttcaagc ggcgagttag
                                                                            480
     ggaacaacaa gattgtgcta tatgtctaga ccggttcaag aagggtgaga ccttggtaca
                                                                            540
     cctaccatgt gcccataagt ttcactccat atgcttattg ccttggctag acaccaatgt
                                                                            600
     ttattgccca tattgtagaa ctgatatttg gaattaaatg ttatattttt ga
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     (2) INFORMATION FOR SEQ ID NO:502:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 196 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
15
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..196
20
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056124
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:502:
     Met Ala Gly Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Phe
                    5
                                         10
     His Gly Gly Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala
25
                                      25
      Ala Ala Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr
                                  40
      Thr Thr Asn His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arg
                              55
30
      Ser Val Arg Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp
                          70
      Gly Ala Ala Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile
                                          90
      Pro Pro Arg Thr Arg Gln Asn Gly Lys Asp Lys Gly Asn Lys Leu Glu
35
                                      105
      Gln Gly Lys Gly Lys Pro Leu Gly Asp Leu Pro Thr Glu Val Val Gly
                                 120
              115
      Leu Lys Lys Ser Arg Gly Arg Leu Met Glu Trp Phe Lys Arg Arg Val
                             135
                                                 140
      Arg Glu Gln Gln Asp Cys Ala Ile Cys Leu Asp Arg Phe Lys Lys Gly
40
                         150
                                             155
      Glu Thr Leu Val His Leu Pro Cys Ala His Lys Phe His Ser Ile Cys
                      165
                                       170
      Leu Leu Pro Trp Leu Asp Thr Asn Val Tyr Cys Pro Tyr Cys Arg Thr
4:5
                 180
                                     185
      Asp Ile Trp Asn
              195
      (2) INFORMATION FOR SEQ ID NO:503:
           (i) SEQUENCE CHARACTERISTICS:
50
                (A) LENGTH: 193 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
55
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..193
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056125
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:503:
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      Met Leu Pro Gly Val Glu Cys Ala Arg Arg Arg Phe His Gly Gly
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Ala Pro Pro Ile Glu Ser Ser Asn Thr Ala Ser Val Ala Ala Ala Ala

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25
     Gly His Val Trp Thr Arg Arg Pro Ser Phe Ser Leu Tyr Thr Thr Asn
                                  40
     His Glu Ser His Gln Ala His Val Ser Phe Ser Glu Arg Ser Val Arg
 5
     Asn Lys Ser Tyr Gly Glu Asp Asn Asp Glu Lys Leu Asp Gly Ala Ala
                                               75
                          70
     Lys Glu Ala Lys Gln Arg Leu Asn Lys Arg Leu Arg Ile Pro Pro Arg
                                           90
10
     Thr Arg Gln Asn Gly Lys Asp Lys Gly Asn Lys Leu Glu Gln Gly Lys
                                      105
      Gly Lys Pro Leu Gly Asp Leu Pro Thr Glu Val Val Gly Leu Lys Lys
                                   120
      Ser Arg Gly Arg Leu Met Glu Trp Phe Lys Arg Arg Val Arg Glu Gln
15
                              135
                                                   140
      Gln Asp Cys Ala Ile Cys Leu Asp Arg Phe Lys Lys Gly Glu Thr Leu
                          150
                                               155
      Val His Leu Pro Cys Ala His Lys Phe His Ser Ile Cys Leu Leu Pro
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                                           170
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      Trp Leu Asp Thr Asn Val Tyr Cys Pro Tyr Cys Arg Thr Asp Ile Trp
                  180
                                       185
      Asn
      (2) INFORMATION FOR SEQ ID NO:504:
25
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 2619 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..2619
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056245
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          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:504:
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                                                                             120
      cacqcqcctc ctqattttca cqcqccqcqc gttccacctc ttatcctctg tcqtqtcqtc
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      ccacttccgg gaaacgactt ggatctagaa aacgacgccg ttttgggtct aactcctcct
                                                                              300
      tettetgacg gtaacggtaa eggtaaagag aaaceggegt etttegetaa aacgttaacg
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      cagtetgacg etaataacgg eggtggtttc teegttecac gttattgege egagacgatt
                                                                              420
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      accaccggtt ggagcacttt cgttaaccag aagaaactaa tcgccggaga ctcaatcgtc
                                                                              660
      ttcctccqtt ctqaatccqq tqacctctqc qtcqqaatcc qccqcqctaa acgcqqcqqt
                                                                              720
      ctcqqatcta acqcaqqatc cqacaatcct taccctqqat tctccqqttt cctccqtqac
                                                                              780
      gacgagtcaa caacaacaac atcgaagcta atgatgatga aacgcaacgg aaacaacgac
      ggaaacgccg cggctacagg gagggttaga gtagaagcag tagcggaagc ggtggcgcgt
                                                                              840
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      gcagcgtgtg gacaagcgtt tgaggttgtt tattatccac gcgctagtac accggagttt
                                                                              900
      tgcgtaaaag cagctgatgt tagatcagca atgaggataa gatggtgtag tggtatgcgt
                                                                              960
      tttaaaatgg cgtttgaaac agaggattct tctagaatca gttggtttat gggtactgtc
                                                                             1020
                                                                             1080
      teegeegtte aagtegetga teeaattegt tggeetaatt caccatggeg teteetteag
      gtagcttggg acgaaccgga tttgttacaa aacgttaagc gggttagtcc gtggttagtc
                                                                             1140
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      aggattccgc agccatttga gtttccattc cacggtacta aattcccgat tttctccccg
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      ggattcgcca acaatggcgg tggcgagtcc atgtgttatc tgtcaaacga caacaataat
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      gctcctgcag gaatacaggg agccaggcaa gctcaacaac tcttcggatc accatctccg
                                                                             1380
      totttgttgt otgatotoaa tottagtagt tacacoggta acaacaagtt acattotoog
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                                                                             1560
      aagaagtetg ttggtteggt taagacteat cagttegtgt tgtteggtea accgatttta
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```

586

									5	986									
	gaga ggtc	aagg attg	tt t ta a	agtg agtt	gctc ttca	g tg t gg	ggtt aatc	aaca tgag	tgg gat	aatt gttg	ata gac	gttt gcac	gcaa actc	gg a ga t	cttg ctct	aggag agacg cggtt agagg	1680 1740 1800 1860		
5	tcgg gacg ggtg ttca	attt aacc gcga aaaa	gt t tt t ca a at t	gact cagt cgtg aatt	catg gatt agaa tttt	t tg t ca a ca t tt	tgta tgaa tttt tgtt	ccgg agca tagt cttt	gat act ttc	gcaa aaac cttc gatg	atg ggc aaa gtg	gtgt taac taat gata	tato aato attt ctat	aa a aa g tt t ac t	cgta gatgg ttct cata	ttgga atatt ttttt tacat	1920 1980 2040 2100		
10	aaac atga ggag caat	ttat agaa tggt ccaa	tt t ac t gg c ct g	gttc catt aaat tgtt	attc caac gagg gctg	t tt t ct g gc c ag	taag cgtt tagg agtg	rttta .cact raaaa rtgac	aaa gtt cca tct	acag atga aaaa atgt	tca tta aac gtg	atataagtaa tataaaaaaa 2160 a attcatttat atcatctatg 2220 a tcttcaccat tcttgtgcta 2280 c aatgtaatga gattctaaag 2340 g tgaagaagag ggggaaagga 2400 a atcattgccc ttaacgattt 2460							
15	tatt tagc taat	attt ctaa aatg INFO	ac a cg a ac a RMAT	ataa ttga atca ION	taag	a ga g tt t ca SEQ	ttta gtct gaag ID N	tgat ctat stttt 10:50	: aaa : caa : tgt )5:	igtag iataa	taa act	aata	atga	at c	gggt	gttta aataa	2460 2520 2580		
20			(A (B (C (D	() LE () TY () ST () TO	NGTH PE: RAND POLC	: 81 amin EDNE GY:	.7 am no ac LSS: line	nino cid ear		is									
25		(ix)	FEA (A (B	TURE  NA  LC  O  O	LE TY L: AME/K DCATI THER CE DE	EY: ON: INFO	pept 18 RMA	ide 317 TION:				_	205	6246	ō				
30	1	Glu	Gln	Glu	Lys 5	Ser	Leu	Asp	Pro	Gln 10	Leu	Trp			15				
0.5			Gly	20	Gln Thr			Ala	25				Asp	30					
35	Pro	Arg 50	35 Val	Pro	Pro	Leu	Ile 55	40 Leu	Cys	Arg	Val	Val 60	45 Ser	Val	Lys	Phe			
40	65				Glu	70					75	=				80			
40					Asn 85 Ser					90					95				
45	Ala	Ser	Phe	100 Ala	Lys	Thr	Leu	Thr 120	105 Gln	Ser	Asp	Ala	Asn 125	110 Asn	Gly	Gly			
	_	130	Ser		Pro	_	135	Cys				140	Phe		_				
50	145	_			Glu Trp	150					155			-	_	160			
				Leu	165 Thr				Ser	170				Gln	175				
55	Leu	Ile	Ala 195	180 Gly	Asp	Ser	Ile	Val 200	185 Phe	Leu	Arg	Ser	Glu 205	190 Ser	Gly	Asp			
		210	Val		Ile		215	Ala				220	Leu						
60	225				Asn Thr	230					235					240			
					245 G1v					250					255				

Gly Asn Asn Asp Gly Asn Ala Ala Ala Thr Gly Arg Val Arg Val Glu

				260					265					270		
	Ala	Val	Ala 275	Glu	Ala	Val	Ala	Arg 280	Ala	Ala	Cys	Gly	Gln 285	Ala	Phe	Glu
5	Val	Val 290	Tyr	Tyr	Pro	Arg	Ala 295	Ser	Thr	Pro	Glu	Phe 300	Cys	Val	Lys	Ala
	Ala 305	Asp	Val	Arg	Ser	Ala 310	Met	Arg	Ile	Arg	Trp 315	Cys	Ser	Gly	Met	Arg 320
	Phe	Lys	Met	Ala	Phe 325	Glu	Thr	Glu	Asp	Ser 330	Ser	Arg	Ile	Ser	Trp 335	Phe
10	Met	Gly	Thr	Val 340	Ser	Ala	Val	Gln	Val 345	Ala	Asp	Pro	Ile	Arg 350	Trp	Pro
			Pro 355					360			_	_	365			
15		370	Asn				375			_		380				
	385		Pro			390					395					400
0.0			Pro		405					410					415	
20			Ser	420	_				425	_	_	_		430		_
	_		Ser 435					440				_	445			
25		450	Ala				455	-				460				
	465		Asn			470					475					480
3.0			Phe		485					490					495	
30			Ser	500					505					510		
			Ala 515				_	520		_			525			
35		530	Gln Asn				535	_				540				
	545		Gly			550					555					560
40			Glu		565					570					575	
40			Thr	580					585					590		
			595 Leu					600					605			
45	_	610					615					620		_		
	625		Pro			630					635					640
50			Asp		645					650					655	
	_		Ile	660	_	_	_		665	_				670		
			675 Ser					680			-		685			
55		690					695					700				
	705		Ile			710					715					720
60			Ser		725					730					735	
			. Phe	740	I				745					750		

		588																
			755					760					765					
	Gly	Lys 770	Pro	Lys	Lys	Gln	Cys 775	Asn	Glu	Ile	Leu	Lys 780	Gln	Ser	Asn	Cys		
5	Val 785	Ala	Ala	Glu	Cys	Asp 790	Ser	Met	Cys	Val	Lys 795	Lys	Arg	Gly	Lys	Gly 800		
		Gly	Tyr	Cys	Ser 805	Pro	Ser	Lys	Lys	Cys 810	Tyr	Cys	Tyr	Tyr	His 815	Cys		
	Pro																	
10	(2)	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 799 amino acids</li><li>(B) TYPE: amino acid</li><li>(C) STRANDEDNESS:</li></ul>																
15		(C) STRANDEDNESS:  (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: peptide  (ix) FEATURE:  (A) NAME/KEY: peptide  (B) LOCATION: 1799																
20									, , ,	`ere	s Sec	T T	) 20°	256247				
20	O (D) OTHER INFORMATION: / Ceres Seq. ID 2056247 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:506:																	
	Met 1							Asn					Tyr	Phe	Ala 15	Gln		
25	Gly	His	Thr	Glu 20	His	Ala	His	Ala	Pro 25	Pro	Asp	Phe	His	Ala 30	Pro	Arg		
	Val	Pro	Pro 35	Leu	Ile	Leu	Cys	Arg 40	Val	Val	Ser	Val	Lys 45	Phe	Leu	Ala		
	Asp	Ala 50	Glu	Thr	Asp	Glu	Val 55	Phe	Ala	Lys	Ile	Thr 60	Leu	Leu	Pro	Leu		
30	Pro 65	Gly	Asn	Asp	Leu	Asp 70	Leu	Glu	Asn	Asp	Ala 75	Val	Leu	Gly	Leu	Thr 80		
	Pro	Pro	Ser	Ser	Asp 85	Gly	Asn	Gly	Asn	Gly 90	Lys	Glu	Lys	Pro	Ala 95	Ser		
35			_	100				Ser	105					110				
			115	_				Glu 120					125					
		130					135	Thr				140	_			_		
40	145			_		150		Ile			155				_	160		
					165			Thr		170					175			
45				180				Leu	185				_	190				
			195					Arg 200					205					
F.O.		210					215	Phe				220						
50	225					230		Leu			235	_				240		
					245			Thr		250					255			
55				260				Ala	265					270				
			275					Pro 280					285					
60		290					295					300		_				
60	305					310		Ser			315					320		
	Thr	val	ser	ΑΙа	val	GΙN	. val	Ala	Asp	Pro	ıте	Arg	Trp	Pro	Asn	ser		

	589															
					325					330					335	
	Pro	Trp	Arg	Leu 340	Leu	Gln	Val	Ala	Trp 345	Asp	Glu	Pro	Asp	Leu 350	Leu	Gln
5	Asn	Val	Lys 355	Arg	Val	Ser	Pro	Trp 360	Leu	Val	Glu	Leu	Val 365	Ser	Asn	Met
	Pro	Thr 370	Ile	His	Leu	Ser	Pro 375	Phe	Ser	Pro	Arg	Lys 380	Lys	Ile	Arg	Ile
	Pro 385	Gln	Pro	Phe	Glu	Phe 390	Pro	Phe	His	Gly	Thr 395	Lys	Phe	Pro	Ile	Phe 400
10			Gly		405					410					415	
			Asp	420					425					430		
15			Gln 435					440					445			
		450	Ser				455					460				
20	465		Ser			470					475					480
20			Asn Val		485					490					495	
			Val	500		_		=	505					510		
25			515 Lys					520					525			
		530	Val				535					540				
30	545		Gly			550					555					560
			Asp		565	_				570			_		575	_
			Glu	580					585	_				590		
35			595 Tyr					600		_			605			
		610	Ser	_	-		615					620				
	625					630					635					640
40			Gly	_	645					650					655	
			Phe	660					665					670		
45			Gly 675					680					685			
		690	Ile	_			695					700	_	_		
50	705		Phe			710					715					720
30			Met		725					730					735	
			Ile Lys	740					745				_	750		
55			755					760					765			
		770					775		_	_		780				атЛ
60	785 (2)		Ser ORMA			790				Cys	795	тАт	пта	Cys	LIO	
	(		) SE	QUEN		HARA	CTER	ISTI	CS:	ds						

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590
             (B) TYPE: amino acid
              (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
5
        (ix) FEATURE:
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..567
             (D) OTHER INFORMATION: / Ceres Seq. ID 2056248
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:507:
10
     Met Met Lys Arg Asn Gly Asn Asp Gly Asn Ala Ala Ala Thr
                                  10
     Gly Arg Val Arg Val Glu Ala Val Ala Glu Ala Val Ala Arg Ala Ala
                                - 25
     Cys Gly Gln Ala Phe Glu Val Val Tyr Tyr Pro Arg Ala Ser Thr Pro
15
                             40
     Glu Phe Cys Val Lys Ala Ala Asp Val Arg Ser Ala Met Arg Ile Arg
     Trp Cys Ser Gly Met Arg Phe Lvs Met Ala Phe Glu Thr Glu Asp Ser
                      70
                                       75
20
     Ser Arg Ile Ser Trp Phe Met Gly Thr Val Ser Ala Val Gln Val Ala
                  85
                                    90
     Asp Pro Ile Arg Trp Pro Asn Ser Pro Trp Arg Leu Leu Gln Val Ala
                    105
               100
     Trp Asp Glu Pro Asp Leu Leu Gln Asn Val Lys Arg Val Ser Pro Trp
25
                   120
     Leu Val Glu Leu Val Ser Asn Met Pro Thr Ile His Leu Ser Pro Phe
                         135
     Ser Pro Arg Lys Lys Ile Arg Ile Pro Gln Pro Phe Glu Phe Pro Phe
                                       155 160
30
     His Gly Thr Lys Phe Pro Ile Phe Ser Pro Gly Phe Ala Asn Asn Gly
                  165
                                    170
     Gly Glu Ser Met Cys Tyr Leu Ser Asn Asp Asn Asn Asn Ala Pro
                                185
     Ala Gly Ile Gln Gly Ala Arg Gln Ala Gln Gln Leu Phe Gly Ser Pro
35
                             200
     Ser Pro Ser Leu Leu Ser Asp Leu Asn Leu Ser Ser Tyr Thr Gly Asn
                                           220
                          215
     Asn Lys Leu His Ser Pro Ala Met Phe Leu Ser Ser Phe Asn Pro Arg
                      230
                                       235
40
     His His His Tyr Gln Ala Arg Asp Ser Glu Asn Ser Asn Asn Ile Ser
                                    250
                   245
     Cys Ser Leu Thr Met Gly Asn Pro Ala Met Val Gln Asp Lys Lys
               260
                                 265
     Ser Val Gly Ser Val Lys Thr His Gln Phe Val Leu Phe Gly Gln Pro
45
                             280
     Ile Leu Thr Glu Gln Gln Val Met Asn Arg Lys Arg Phe Leu Glu Glu
                         295
                                           300
     Glu Ala Glu Ala Glu Glu Lys Gly Leu Val Ala Arg Gly Leu Thr
                                       315
                      310
50
     Trp Asn Tyr Ser Leu Gln Gly Leu Glu Thr Gly His Cys Lys Val Phe
                   325
                                    330
     Met Glu Ser Glu Asp Val Gly Arg Thr Leu Asp Leu Ser Val Ile Gly
                340
                                 345
     Ser Tyr Gln Glu Leu Tyr Arg Lys Leu Ala Glu Met Phe His Ile Glu
55
                             360
     Glu Arg Ser Asp Leu Leu Thr His Val Val Tyr Arg Asp Ala Asn Gly
                          375
                                            380
     Val Ile Lys Arg Ile Gly Asp Glu Pro Phe Ser Asp Phe Met Lys Ala
                                        395 400
                      390
     Thr Lys Arg Leu Thr Ile Lys Met Asp Ile Gly Gly Asp Asn Val Arg
60
                   405
                                  410
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Asn Ile Phe Ser Phe Leu Gln Ile Ile Phe Phe Phe Phe Phe Lys

591 425 420 430 Lys Leu Ile Phe Phe Leu Phe Phe Ser Asp Gly Gly Tyr Tyr Thr His 440 445 Ile His Ile Tyr Ser Ser Asp Tyr Ile Tyr Ile Tyr Ile Tyr Ile Tyr 5 455 Ile Ser Asn Ile Lys Lys Lys Leu Ile Leu Phe Ile Leu Leu Ser Leu 470 475 Lys Thr Val Asn Ser Phe Ile Ser Ser Met Met Lys Lys Leu Ile Gln 485 490 10 Leu Ser Phe Thr Val Met Ile Ile Phe Thr Ile Leu Val Leu Gly Val 500 505 Val Ala Asn Glu Gly Leu Gly Lys Pro Lys Lys Gln Cys Asn Glu Ile 520 525 Leu Lys Gln Ser Asn Cys Val Ala Ala Glu Cys Asp Ser Met Cys Val 15 535 540 530 Lys Lys Arg Gly Lys Gly Ala Gly Tyr Cys Ser Pro Ser Lys Lys Cys 550 555 Tyr Cys Tyr Tyr His Cys Pro 565 20 (2) INFORMATION FOR SEQ ID NO:508: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 776 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 25 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..776 30 (D) OTHER INFORMATION: / Ceres Seq. ID 1941142 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:508: 60 agtogoctoc ttootgogoc googoogoog cogocgoogo cgctaccaco gootogogo 120 cgcctgcggc tccgcgtttc cgcacggaag ttggtgaagg tttcttgagg ctgagaacat ggtgctgaag acggaacttt gccgcttcag cggccagaag atttatcctg ggaaaggcat 180 35 tagatttatc cgtgctgatt ctcaggtctt cctttttgcc aactcgaaat gcaagcgcta 240 cttccacaac cgcctgaagc ctgcaaagct tacctggaca gcaatgtaca ggaagcagca 300 caagaaggat atccatgctg aagcggtaaa gaagaggcgc cgcgccacca agaagccata 360 420 ctccaggtca attgtgggtg cttccttgga agtaatccag aagaagagag ctgagaagcc agaggtccgc gatgctgcta gagaggctgc tcttcgtgag atcaaggagc gcatcaagaa 480 40 gaccaaggat gagaagaaag cgaagaaggc ggaggtgagc aagtcccaga agacgcagac 540 aaaqqqtqcq qtccaqaaqq qttccaaqqq ccccaaqttq qqcqqcqqtg qtqqqaaqcq 600 660 ctgaaagaac ttagtgtcgt ttctcgacat tgcagtcgtt ccttagccaa agccactttc 720 gtagaacctg tgttgaattt gcaagactta ttcaagcgtt gcttgtgcgt gctaaatacc atggcaagag aacggattta tatttatgcc tgaaaaaaat gaccgttcat attctt 45 (2) INFORMATION FOR SEQ ID NO:509: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 161 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 50 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..161 55 (D) OTHER INFORMATION: / Ceres Seq. ID 1941143 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:509: Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr 10 Pro Gly Lys Gly Ile Arg Phe Ile Arg Ala Asp Ser Gln Val Phe Leu 60 25

Phe Ala Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Arg Leu Lys Pro 35 40 45

592 Ala Lys Leu Thr Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp 5.5 Ile His Ala Glu Ala Val Lys Lys Arg Arg Ala Thr Lys Lys Pro 70 75 5 Tyr Ser Arg Ser Ile Val Gly Ala Ser Leu Glu Val Ile Gln Lys Lys 90 85 Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu 100 105 Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala 10 120 Lys Lys Ala Glu Val Ser Lys Ser Gln Lys Thr Gln Thr Lys Gly Ala 135 Val Gln Lys Gly Ser Lys Gly Pro Lys Leu Gly Gly Gly Gly Lys 15 Arq (2) INFORMATION FOR SEQ ID NO:510: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 106 amino acids 20 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 25 (A) NAME/KEY: peptide (B) LOCATION: 1..106 (D) OTHER INFORMATION: / Ceres Seq. ID 1941144 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:510: Met Tyr Arg Lys Gln His Lys Lys Asp Ile His Ala Glu Ala Val Lys 30 10 Lys Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly 25 Ala Ser Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val 40 35 Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile 55 Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Ala Glu Val Ser Lys 75 Ser Gln Lys Thr Gln Thr Lys Gly Ala Val Gln Lys Gly Ser Lys Gly 40 85 Pro Lys Leu Gly Gly Gly Gly Lys Arg 100 (2) INFORMATION FOR SEQ ID NO:511: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 1024 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 50 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..1024 (D) OTHER INFORMATION: / Ceres Seq. ID 1942975 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:511: 55 ataaattcct ccgctcccct tcgccgcttc ctgtcgcgcc gccgctcccc tcccaaccca 60 120 ctageteaaa accetagage tetegeetge tegeeteege egeeggageg ageteeecat 180 cttcatccgt caaccatggc gacccagatc agcaagaaga agaagttcgt cagcgacggt gttttctacg ccgagctcaa tgagatgctg acgcgggagc tggcggagga cggctactct 240 ggcgtggagg tgcgcgtcac gccgatgcqc acqqagatca tcatccqcqc cacqcqcacg 300 60 cagaacgtgc tcggcgagaa gggccgtagg atcagggagc tcacctccgt cgtccagaag 360 420 aggttcaact teeetqagaa tggcqttqaq etttatgeeg agaaggtegt caaccqtgqq 480 ctctgtgcca tcgcgcaggc cgagtccctc cgttacaagc tcctcggtgg ccttgccgtc

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									;	593							
																gtgag	540
	gtga	tcgt	ga ç	gtgga	aagc	t ca	gggc	ccaa	aga	igcca	agt	ccat	gaag	gtt c	caago	gatggc	600
		_					_	_			_					cacgtt	660
_																ccgaag	720
5																aggac	780
																gtgtc	840
	cttg	gaaat	gg c	cagtt	tgat	c gg	gtcc	taga	ı taç	gttt	.cta	tctt	acga	ata a	aggac	caatgc	900
	taag	gcaç	gtt g	ggtta	acctt	a at	cttt	atgo	, aca	gato	rcta	tgt	gttgc	ctt d	ctgtc	gcact	960
	tttg	ıttta	ag c	ctaac	gtttt	g cg	gaga	caac	, ata	actta	tat	atgt	tccc	cga t	gaaa	ittatg	1020
10	gttc	;															
	(2)			NOI													
		(i)	SEÇ	QUENC	CE CH	IARAC	TERI	STIC	CS:								
			( Z	A) LE	ENGTH	I: 27	'4 am	nino	acio	ds							
			( E	3) TY	PE:	amin	o ac	cid									
15			( (	C) SI	RANI	DEDNE	SS:										
	(D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide																
		(ix)		ATURE													
				A) NA													
20				3) LO													
(D) OTHER INFORMATION: / Ceres Seq. ID 1942976 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:512:																	
	Ile	Asn	Ser	Ser	Ala	Pro	Leu	Arg	Arg		Leu	Ser	Arg	Arg	_	Ser	
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25	Pro	Pro	Asn	Pro	Leu	Ala	Gln	Asn		Arg	Ala	Leu	Ala	-	Ser	Pro	
	_		_	20	_		_		25		_			30	_		
	Pro	Pro		Glu	Arg	Ala	Pro		Leu	His	Pro	Ser		Met	Ala	Thr	
	~ >		35	_	_	_	_	40		_	_		45	_,	_		
20	GIn		Ser	Lys	ГÀЗ	Lys		Phe	Val	Ser	Asp	_	Val	Phe	Tyr	Ala	
30		50	_			_	55	_		_		60	_	- 3	_		
		Leu	Asn	Glu	Met		Thr	Arg	Glu	Leu		GLu	Asp	GLy	Tyr		
	65		<b>a</b> 1		_	70	m.	_		-	75	<b>.</b>		- 1	- 1	80	
	GTĀ	Va⊥	Glu	Val	_	٧al	Thr	Pro	Met	_	Thr	GLu	IТе	тте		Arg	
25	7.1.	m \	70	m 1	85	7)	77± 1	T	C1	90	T	C1	7)	7)	95	7)	
35	АТа	Thr	Arg	Thr	GIN	ASI	vаı	Leu	_	GIU	глг	GΤΔ	Arg		тте	Arg	
	Clu	T 0.11	mb w	100	77-1	17-1	C1 n	T	105	Dha	7.00	Dh =	Dwa	110	7.00	C1	
	GIU	Leu	115	Ser	٧aı	∨а⊥	Gill	120	Arg	Pne	ASII	Pne	125	GLU	ASII	GIY	
	57 n 1	Cin		Tyr	71-	Clu	T		77-1	Λαρ	λκα	C1		Cuc	77.70	T l o	
40	Val	130	ьеи	ТЭТ	Ата	GIU	135	۷ат	vaı	ASII	AIG	140	теп	Cys	Ата	TTG	
40	חות		ת ז ת	Glu	Sor	T ON		Тих	Tuc	Tou	Τ ο ι ι		C1.,	T OU	7.1.	7751	
	145	GIII	Ата	GIU	ser	150	Arg	тут	гуѕ	ьеи		дту	GTA	ьеч	Ата	160	
		7) ra	Λla	Cys	Тих		U = 1	Lou	λκα	Тих	155	Mot	Glu	Sor	Glv		
	ALG	лту	Ата	Cys	165	Эту	var	пеа	Arg	170	vaı	rie c	GIU	Jer	175	AIA	
45	Lve	C1 v	Cve	Glu		Tlo	Val	Sar	Glv		LAu	Ara	ΔΙα	Gln		Δ1 =	
40	пуз	GIY	Cys	180	vaı	116	vaı	Ser	185	гуз	ьeu	ALG	Ата	190	Arg	AIG	
	Lus	Sar	Met	Lys	Phe	T.vs	Asn	G1 v		Mot	Τlြ	Sar	Sar		Gln	Pro	
	пур	261	195	цуз	1110	Бүр	115P	200	гуг	PICC	110	Ser	205	ary	1110	110	
	Val	Aen		Tyr	Tla	Δsn	Ser	_	Val	Δrα	His	U=1		Len	Δrα	G1n	
50	val	210	GIU	тАт	116	nsp	215	лта	Val	Arg	1113	220	цец	пеа	Arg	GIII	
<b>J</b> 0	Glv		T.eu	Gly	Tle	1.00		T.VS	Tle	Met	T.em		Trn	Asn	Pro	T.Ve	
	225	Val	пса	OTY	110	230	٠٠٠	дуо	110	1100	235	пор	115	тэр	110	240	
		I.ve	Val	Gly	Pro		Thr	Pro	T.e.i	Pro		T.e.i	Val	Thr	Tlo		
	- T Y	دړب	v u i	⊸ ± y	245		- 444	0	<u> </u>	250	ر ين د	<u> </u>	· u ⊥	* * * * *	255		
55	Thr	Pro	Lvs	Asp		Asp	Glu	Pro	Ara		Pro	Val	Len	Ala		Pro	
<b>~ ~</b>	-114		-,5	260	<u> </u>				265		0			270	0		
	G] 11	Val		200					_ 55								
	(2)	TNE	ORMA	тт∩м	FOR	SEO	TD.	NO • 5	13.								

- (2) INFORMATION FOR SEQ ID NO:513:
- (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 229 amino acids
- (B) TYPE: amino acid

594

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(C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
 5
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..229
               (D) OTHER INFORMATION: / Ceres Seq. ID 1942977
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:513:
     Met Ala Thr Gln Ile Ser Lys Lys Lys Phe Val Ser Asp Gly Val
10
                                         10
     Phe Tyr Ala Glu Leu Asn Glu Met Leu Thr Arg Glu Leu Ala Glu Asp
                 20
                                     25
     Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile
                                 40
15
      Ile Ile Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg
                             55
                                                 60
     Arg Ile Arg Glu Leu Thr Ser Val Val Gln Lys Arg Phe Asn Phe Pro
                         70
                                             75
     Glu Asn Gly Val Glu Leu Tyr Ala Glu Lys Val Val Asn Arg Gly Leu
20
                     85
                                         90
      Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly
                  100
                                     105
      Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Tyr Val Met Glu
                                 120
                                                     125
25
      Ser Gly Ala Lys Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala
                             135
                                                 140
      Gln Arg Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Ile Ser Ser
                                            155
                         150
     Gly Gln Pro Val Asn Glu Tyr Ile Asp Ser Ala Val Arg His Val Leu
30
                     165
                                         170 175
     Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Leu Asp Trp
                                     185
                 180
      Asp Pro Lys Gly Lys Val Gly Pro Ile Thr Pro Leu Pro Asp Leu Val
             195
                                200
                                          205
35
      Thr Ile His Thr Pro Lys Asp Glu Asp Glu Pro Arg Pro Pro Val Leu
         210
                             215
      Ala Pro Pro Glu Val
      (2) INFORMATION FOR SEQ ID NO:514:
40 .
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 206 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..206
                (D) OTHER INFORMATION: / Ceres Seq. ID 1942978
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:514:
      Met Leu Thr Arg Glu Leu Ala Glu Asp Gly Tyr Ser Gly Val Glu Val
                    5
                                         10
      Arg Val Thr Pro Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr
                                     25
55
      Gln Asn Val Leu Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser
                                 40
      Val Val Gln Lys Arg Phe Asn Phe Pro Glu Asn Gly Val Glu Leu Tyr
      Ala Glu Lys Val Val Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu
60
                                             75
      Ser Leu Arg Tyr Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys
```

90

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595
     Tyr Gly Val Leu Arg Tyr Val Met Glu Ser Gly Ala Lys Gly Cys Glu
                 100
                                    105
     Val Ile Val Ser Gly Lys Leu Arg Ala Gln Arg Ala Lys Ser Met Lys
                                 120
                                                    125
 5
     Phe Lys Asp Gly Tyr Met Ile Ser Ser Gly Gln Pro Val Asn Glu Tyr
                             135
     Ile Asp Ser Ala Val Arg His Val Leu Leu Arg Gln Gly Val Leu Gly
                         150
                                            155
     Ile Lys Val Lys Ile Met Leu Asp Trp Asp Pro Lys Gly Lys Val Gly
10
                     165
                                         170
     Pro Ile Thr Pro Leu Pro Asp Leu Val Thr Ile His Thr Pro Lys Asp
                 180
                      185
     Glu Asp Glu Pro Arg Pro Pro Val Leu Ala Pro Pro Glu Val
             195
                                 200
15
      (2) INFORMATION FOR SEQ ID NO:515:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1069 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
20
                (D) TOPOLOGY: linear
          (11) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
               (B) LOCATION: 1..1069
25
                (D) OTHER INFORMATION: / Ceres Seq. ID 1944349
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:515:
     acttegttte tgcetttgae eagetettge geteggetta tetegtgegg ttgttgetag
                                                                           60
      aaccgtccct aattccattc ccgtgatctg gtttagtcat ttcatggttt tatttgttgc
                                                                          120
      atctgtggtg aatctgtggt cacttagatc aacataggaa gatgtacgcc cagttgagtt
                                                                          180
30
     ttgagtcggc tgttggtgtt tccctaacta gggactgagg gtctgaagct ctatacctgt
                                                                          240
      tgcaatcgcg tagttgtgcg tgctagatgt atcagagaat ccctttgagt agaaagcaca
                                                                          300
                                                                          360
      cgcacacage tgccacggta ttggcattcg taggtatctg tttcactgac gtgctacctg
                                                                          420
      cacgaggata gcacctgata tttgtgcgct ttttttggtg aagatttttt gaagctgaga
                                                                          480
      agatggtgct gaagacggaa ctctgccgct tcagtggcca gaagatctat cctgggaaag
35
                                                                          540
     gcatccggtt tattcgtgct gattctcagg ttttcctttt tgccaactct aaatgcaagc
      gctacttcca caaccgcctg aagcctgcaa agcttacctg gacagcaatg tacaggaagc
                                                                          600
      660
                                                                          720
      catactcgag gtccattgtg ggtgcttcct tggaagtgat ccagaagaag agagctgaga
                                                                          780
      agectgaggt ccgtgatgct gctagagaag ctgctcttcg tgagatcaag gagcgcatca
40
                                                                          840
      agaagaccaa ggacgagaag aaggcgaaga aggctgaggt gtccaagtcc cagaaggcgc
                                                                          900
      agacqaaqqq cqctqtccaq aaqqqttcca aqqqccccaa qttqqqcqqc qqcqqtqqca
                                                                         960
      aacgctgaaa aggaacgaag tgtcatttcg agacctcacc aaagtcacct ttgtagtacc
      tgggttaatt ttgcattttc gttcagagtt acttgtgact cttaactacc atgtcaagag
                                                                         1020
      aatggattcg ttattttat atattgggag aaatgatcgc cgttccttc
45
      (2) INFORMATION FOR SEQ ID NO:516:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 161 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
50
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..161
55
                (D) OTHER INFORMATION: / Ceres Seq. ID 1944350
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:516:
      Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr
                                         10
      Pro Gly Lys Gly Ile Arg Phe Ile Arg Ala Asp Ser Gln Val Phe Leu
60
                                  25
                 20
      Phe Ala Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Arg Leu Lys Pro
                                 40
```

```
596
     Ala Lys Leu Thr Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp
                            5.5
     Ile His Ala Glu Ala Val Lys Lys Arg Arg Ala Thr Lys Lys Pro
                        70
                                        75
 5
     Tyr Ser Arg Ser Ile Val Gly Ala Ser Leu Glu Val Ile Gln Lys Lys
                    85
                                       90
     Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu
                                   105
     Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala
10
                               120
     Lys Lys Ala Glu Val Ser Lys Ser Gln Lys Ala Gln Thr Lys Gly Ala
         130 135
                                               140
     Val Gln Lys Gly Ser Lys Gly Pro Lys Leu Gly Gly Gly Gly Lys
                        150
                                           155
15
     Arg
      (2) INFORMATION FOR SEQ ID NO:517:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 106 amino acids
20
               (B) TYPE: amine acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
25
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..106
               (D) OTHER INFORMATION: / Ceres Seq. ID 1944351
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:517:
     Met Tyr Arg Lys Gln His Lys Lys Asp Ile His Ala Glu Ala Val Lys
30
                                       10
     Lys Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
               20
                                25
     Ala Ser Leu Glu Val Ile Gln Lys L's Arg Ala Glu Lys Pro Glu Val
                               40
35
     Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
                           55
     Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Ala Glu Val Ser Lys
                        70
                                   75
     Ser Gln Lys Ala Gln Thr Lys Gly Ala Val Gln Lys Gly Ser Lys Gly
40
                 85
      Pro Lys Leu Gly Gly Gly Gly Lys Arg
                100
      (2) INFORMATION FOR SEQ ID NO:518:
          (i) SEQUENCE CHARACTERISTICS:
45
               (A) LENGTH: 89 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
50
         (ix) FEATURE:
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..89
               (D) OTHER INFORMATION: / Ceres Seq. ID 1944352
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:518:
55
     Met Leu Leu Glu Lys Leu Phe Val Arg Ser Arg Ser Ala Ser Arg
                                       10
     Arg Pro Arg Thr Arg Arg Arg Arg Arg Leu Arg Cys Pro Ser Pro
                                    25
      Arg Arg Arg Arg Arg Ala Leu Ser Arg Arg Val Pro Arg Ala Pro
60
                             40
      Ser Trp Ala Ala Val Ala Asn Ala Glu Lys Glu Arg Ser Val Ile
```

597 Ser Arg Pro His Gln Ser His Leu Cys Ser Thr Trp Val Asn Phe Ala 70 Phe Ser Phe Arg Val Thr Cys Asp Ser 85 (2) INFORMATION FOR SEQ ID NO:519: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1035 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..1035 15 (D) OTHER INFORMATION: / Ceres Seq. ID 1964011 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:519: aacatataaa ttacqtqctc cctcttcqcc qccqtctctc qcqccqccac ttccctcccq 60 acceaetege ccaaaacegt agegeeeteg cetgetegee geegeegteg ttgegagete 120 cccatcttcc tccqttaacc atqqcqaccc aqatcaqcaa qaaqaagaag ttcqtcaqcq 180 20 acqqcqtqtt ctacqccqaq ctcaatqaqa tqctqacqcq qqaqctcqcq qaqqacqqct 240 actooggogt ggaggtgogt gtoacacoga tgogcaogga gatoatcato cgogcaaogc 300 GCacqcaqaa cqtqctcqqc qaqaaqqqcc qcaqGatcaq qqaqctcacq tctqtqqtcc 360 agaagaggtt caacttcccc gagaacggcg tcgagctcta cgccgagaag gtcgtcaacc 420 degreetetg caccategeg caggetgagt eccteegeta caageteete ggaggeettg 480 25 ccgtccggag ggcctgctat ggtgttcttc gctttgttat ggagagtggt gccgagggtt 540 600 gtgaggtcat tgtgagtgga aagctcaggg ctcaaagagc caagtccatg aagttcaagg atggctacat gatctcatct ggtcagccag tgaatgagta catcgactca gctgtgagac 660 720 acgtectect cagacagggt gttetgggta teaaggtgaa gateatgett gaetgggaee 780 cgaagggcaa ggttggcccg atcacccctc ttccggacct ggtgaccatc cataccccga 30 aggaggagga cgagctgcgc cctcctgtct tggcagctga ggtctaaggt cccaacagcg 840 tcctcgaaat ggcagtttga tcgggtccta gctaggtttg atcttatgat aagggcagtg 900 ctaagacagt tcgttccgtg tcacttctgt ttaaaattaa gttttgccga gacaagctat 960 atgtaccgga tgaaagtgtg gttcc+tatt gtatgctctg gaattttatt tttggtatgt 1020 ttatttttt gccct 35 (2) INFORMATION FOR SEQ ID NO:520: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 274 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 40 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..274 45 (D) OTHER INFORMATION: / Ceres Seq. ID 1964012 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:520: His Ile Asn Tyr Val Leu Pro Leu Arg Arg Leu Ser Arg Arg His 10 Phe Pro Pro Asp Pro Leu Ala Gln Asn Arg Ser Ala Leu Ala Cys Ser 50 20 25 Pro Pro Pro Ser Leu Arg Ala Pro His Leu Pro Pro Leu Thr Met Ala 40 45 Thr Gln Ile Ser Lys Lys Lys Phe Val Ser Asp Gly Val Phe Tyr 55 60 55 Ala Glu Leu Asn Glu Met Leu Thr Arg Glu Leu Ala Glu Asp Gly Tyr 75 70 Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile Ile Ile 90 Arg Ala Thr Arg Thr Gln Asn Val Leu Gly Glu Lys Gly Arg Arg Ile 60 100 105 110 Arg Glu Leu Thr Ser Val Val Gln Lys Arg Phe Asn Phe Pro Glu Asn

120

125

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598
     Gly Val Glu Leu Tyr Ala Glu Lys Val Val Asn Arg Gly Leu Cys Ala
                       135 140
     Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly Leu Ala
                           155
                      150
 5
     Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu Ser Gly
                  165
                        170 175
     Ala Glu Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala Gln Arg
              180 185 190
     Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Ile Ser Ser Gly Gln
10
      195 200 205
     Pro Val Asn Glu Tyr Ile Asp Ser Ala Val Arg His Val Leu Leu Arg
                         215
                                           220
     Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Leu Asp Trp Asp Pro
                     230
                                        235
15
     Lys Gly Lys Val Gly Pro Ile Thr Pro Leu Pro Asp Leu Val Thr Ile
                  245
                                    250
     His Thr Pro Lys Glu Glu Asp Glu Leu Arg Pro Pro Val Leu Ala Ala
                                 265
     Glu Val
20
     (2) INFORMATION FOR SEQ ID NO:521:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 228 amino acids
              (B) TYPE: amino acid
25
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
30
              (B) LOCATION: 1..228
              (D) OTHER INFORMATION: / Ceres Seq. ID 1964013
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:521:
     Met Ala Thr Gln Ile Ser Lys Lys Lys Phe Val Ser Asp Gly Val
                                     10
35
     Phe Tyr Ala Glu Leu Asn Glu Met Leu Thr Arq Glu Leu Ala Glu Asp
               20
                       25
     Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Met Arg Thr Glu Ile
                             40
     Ile Ile Arq Ala Thr Arq Thr Gln Asn Val Leu Gly Glu Lys Gly Arq
40
     Arg Ile Arg Glu Leu Thr Ser Val Val Gln Lys Arg Phe Asn Phe Pro
                                     75
     Glu Asn Gly Val Glu Leu Tyr Ala Glu Lys Val Val Asn Arg Gly Leu
                                     90
45
     Cys Ala Ile Ala Gln Ala Glu Ser Leu Arg Tyr Lys Leu Leu Gly Gly
                                 105
                                                  110
     Leu Ala Val Arg Arg Ala Cys Tyr Gly Val Leu Arg Phe Val Met Glu
                                               125
                              120
     Ser Gly Ala Glu Gly Cys Glu Val Ile Val Ser Gly Lys Leu Arg Ala
50
                                           140
                         135
     Gln Arg Ala Lys Ser Met Lys Phe Lys Asp Gly Tyr Met Ile Ser Ser
                      150 155 160
     Gly Gln Pro Val Asn Glu Tyr Ile Asp Ser Ala Val Arg His Val Leu
                  165
                                    170 175
55
     Leu Arg Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Leu Asp Trp
               180
                                185
     Asp Pro Lys Gly Lys Val Gly Pro Ile Thr Pro Leu Pro Asp Leu Val
                            200 205
     Thr Ile His Thr Pro Lys Glu Glu Asp Glu Leu Arg Pro Pro Val Leu
60
                         215
                                           220
     Ala Ala Glu Val
     225
```

599

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(2) INFORMATION FOR SEO ID NO:522:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 205 amino acids
                (B) TYPE: amino acid
 5
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
10
                (B) LOCATION: 1..205
                (D) OTHER INFORMATION: / Ceres Seq. ID 1964014
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:522:
      Met Leu Thr Arg Glu Leu Ala Glu Asp Gly Tyr Ser Gly Val Glu Val
                      5
                                          10
15
      Arg Val Thr Pro Met Arg Thr Glu Ile Ile Ile Arg Ala Thr Arg Thr
                                      25
      Gln Asn Val Leu Gly Glu Lys Gly Arg Arg Ile Arg Glu Leu Thr Ser
                                  40
      Val Val Gln Lys Arg Phe Asn Phe Pro Glu Asn Gly Val Glu Leu Tyr
20
                              55
      Ala Glu Lys Val Val Asn Arg Gly Leu Cys Ala Ile Ala Gln Ala Glu
                          7.0
      Ser Leu Arg Tyr Lys Leu Leu Gly Gly Leu Ala Val Arg Arg Ala Cys
                      85
                                           90
25
      Tyr Gly Val Leu Arg Phe Val Met Glu Ser Gly Ala Glu Gly Cys Glu
                  100
                                      105
                                                           110
      Val Ile Val Ser Gly Lys Leu Arg Ala Gln Arg Ala Lys Ser Met Lys
              115
                                  120
      Phe Lys Asp Gly Tyr Met Ile Ser Ser Gly Gln Pro Val Asn Glu Tyr
30
          130
                              135
                                                   140
      Ile Asp Ser Ala Val Arg His Val Leu Leu Arg Gln Gly Val Leu Gly
                          150
                                               155
      Ile Lys Val Lys Ile Met Leu Asp Trp Asp Pro Lys Gl Lys Val Gly
                      165
                                           170
35
      Pro Ile Thr Pro Leu Pro Asp Leu Val Thr Ile His Thr Pro Lys Glu
                  180
                                      185
      Glu Asp Glu Leu Arg Pro Pro Val Leu Ala Ala Glu Val
                                  200
      (2) INFORMATION FOR SEQ ID NO:523:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 818 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..818
                (D) OTHER INFORMATION: / Ceres Seq. ID 1983854
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:523:
      gatataagaa accctagcgg agccactaac acagtcgcct ccttcctgcr ccgccgccgc
                                                                             60
      cgccgccgcc gccgctacca ccgcctcgcg gccgcctgcg gctccgcgtt tccgcacgga
                                                                             120
                                                                             180
      agttggtgaa ggtttcttga ggctgagaac atggtgctga agacggaact ttgccgcttc
      agcgscagaa gatttatcct gggaaaggca ttagatttat ccgtgctgat tctcaggtct
                                                                             240
55
      tcctttttgc caactcgaaa tgcaagcgct acttccacaa ccgcctgaag cctgcaaagc
                                                                             300
                                                                             360
      ttacctggac agcaatgtac aggaagcagc acaagaagga tatccatgct gaagcggtaa
                                                                             420
      agaagaggcg ccgcgccacc aagaagccat actccaggtc aattgtgggt gcttccttgg
      aagtaatcca gaagaagaga gctgagaagc cagaggtccg cgatgctgct agagaagctg
                                                                             480
      ctcttcgtga gatcaaggag cgcatcaaga agaccaagga tgagaagaaa gcgaagaagg
                                                                             540
60
      cggaggtgag caagtcccag aagacgcaga caaagggtgc ggtccagaag ggttccaagg
                                                                            600
                                                                             660
      gccccaagtt gggcggcggt ggtgggaagc gctgaaagaa cttagtgtcg tttctcgaca
```

ttgcagtcgt tccttagcca aagccacttt cgtagaacct gtgttgaatt tgcaagactt

```
attcaagegt tgcttgtgeg tgctaaatac catggcaaga gaacggattt atatttatge
                                                                             780
      ctgaaaaaa tgaccgttca tattcttaac tcatcttg
      (2) INFORMATION FOR SEQ ID NO:524:
           (i) SEQUENCE CHARACTERISTICS:
 5
                (A) LENGTH: 106 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
10
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..106
                (D) OTHER INFORMATION: / Ceres Seq. ID 1983855
          (xi; SEQUENCE DESCRIPTION: SEQ ID NO:524:
15
      Met Tyr Arg Lys Gln His Lys Lys Asp Ile His Ala Glu Ala Val Lys
                      5
                                          10
      Lys Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
                  20
                                      25
      Ala Ser Leu Glu Val Ile Gln Lvs Lys Arg Ala Glu Lys Pro Glu Val
20
              35
                                  40
                                                       45
      Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
                              55
          50
                                                   60
      Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Ala Glu Val Ser Lys
                                               75
                          70
      65
25
      Ser Gln Lys Thr Gln Thr Lys Gly Ala Val Gln Lys Gly Ser Lys Gly
                                           90
                      85
      Pro Lys Leu Gly Gly Gly Gly Lys Arg
                  100
      (2) INFORMATION FOR SEQ ID NO:525:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 802 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..802
                (D) OTHER INFORMATION: / Ceres Seq. ID 1990261
40
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:525:
      gatataagaa accctagcgg mscactaaya cagtcgcctc cttcctgcgc cgccgccgc
                                                                              60
      geogeogeta ceaeegeete geggeegeet geggeteege gttteegeae ggaagttggt
                                                                             120
                                                                             180
      gaaggtttct tgaggctgag aacatggtgc tgaagacgga actttgccgc ttcagcggcc
                                                                             240
      agaagattta tootgggaaa ggoattagat ttatoogtgo tgattotoag gtottoottt
45
      ttgccaactc gaaatgcaag cgctacttcc acaaccgcct gaagcctgca aagcttacct
                                                                             300
      ggacagcaat gtacaggaag cagcacaaga aggatatcca tgctgaagcg gtaaagaaga
                                                                             360
      ggcgccgcgc caccaagaag ccatactcca ggtcaattgt gggtgcttcc ttggaagtaa
                                                                             420
      tccagaagaa gagagctgag aagccagagg tccgcgatgc tgctagagaa gctgctcttc
                                                                             480
      gtgagatcaa ggagcgcatc aagaagacca aggatgagaa gaaagcgaag aaggcggagg
                                                                             540
50
                                                                             600
      tgagcaagtc ccagaagacg cagacaaagg gtgcggtcca gaagggttcc aagggcccca
                                                                             660
      agttgggcgg cggtggtggg aagcgctgaa agaacttagt gtcgtttctc gacattgcag
      tcgttcctta gccaaagcca ctttcgtaga acctgtgttg aatttgcaag acttattcaa
                                                                             720
                                                                             780
      gcgttgcttg tgcgtgctaa ataccatggc aaqagaacgg atttatattt atgcctgaaa
      aaaaatgacc gttcatattc tt
55
      (2) INFORMATION FOR SEQ ID NO:526:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 161 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
60
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
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601

```
(A) NAME/KEY: peptide
               (B) LOCATION: 1..161
               (D) OTHER INFORMATION: / Ceres Seq. ID 2061972
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:526:
 5
     Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr
                                         10
     Pro Gly Lys Gly Ile Arg Phe Ile Arg Ala Asp Ser Gln Val Phe Leu
                                    25
      Phe Ala Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Arg Leu Lys Pro
10
     Ala Lys Leu Thr Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp
      Ile His Ala Glu Ala Val Lys Lys Arg Arg Arg Ala Thr Lys Lys Pro
                         70
                                             75
15
     Tyr Ser Arg Ser Ile Val Gly Ala Ser Leu Glu Val Ile Gln Lys Lys
                     85
                                         90
     Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu
                 100
                                     105
     Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala
20
                                 120
      Lys Lys Ala Glu Val Ser Lys Ser Gln Lys Thr Gln Thr Lys Gly Ala
       130
                             135
     Val Gln Lys Gly Ser Lys Gly Pro Lys Leu Gly Gly Gly Gly Lys
     145
                         150
                                             155
25
     Arq
      (2) INFORMATION FOR SEQ ID NO:527:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 106 amino acids
30
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
35
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..106
               (D) OTHER INFORMATION: / Ceres Seq. ID 2061973
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:527:
     Met Tyr Arg Lys Gln His Lys Lys Asp Ile His Ala Glu Ala Val Lys
40
                                        10
     Lys Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly
                                 25
     Ala Ser Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val
                                 40
45
      Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile
                             55
      Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Ala Glu Val Ser Lys
                         7.0
                                            75
      Ser Gln Lys Thr Gln Thr Lys Gly Ala Val Gln Lys Gly Ser Lys Gly
50
                    85
      Pro Lys Leu Gly Gly Gly Gly Lys Arg
      SEQ TABLE 2
55
      (2) INFORMATION FOR SEQ ID NO:1:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 494 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
60
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
```

(ix) FEATURE:

```
602
                (A) NAME/KEY: -
                (B) LOCATION: 1..494
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007802
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
 5
      aacaatcgga agaaaggaga agacgaagat gtcgttggta tggttggaag cgatgttgcc
     tctcggaatc atcggtggga tgctctgtat catgggcaat tctcagtact acatccacaa
      agcttatcat ggccgtccta agcacatcgg ccacgatgaa tgggatgttg ctatggaaag
      acgcgacaag aaagtcgtcg agaaagctgc agctccttcc tcatgattcg ctttatctct
      tttgtgttcc traggggctt aaggtgacca cttgtgggtga caaataaagt gcattccaga
                                                                            300
10
     agaagaagaa gctgggggat ctagtacttt cattcccatt tgattttcct tggacatatt
                                                                            360
     aaagetttea qaaateagae etkaataaca tttggtttat caatatttet etattegtga
                                                                            420
      tattttgtat gettttaagt tgeatgatae actgaactae tetettgetg tgtgagtgaa
                                                                            480
      taaatgaata toto
      (2) INFORMATION FOR SEQ ID NO:2:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 94 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                .D) TOPOLOGY: linear
20
          ii) MOLECULE TYPE: peptide
          ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..94
                ID) OTHER INFORMATION: / Ceres Seq. ID 1007803
25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
      Gln Ser Glu Glu Arg Arg Arg Arg Cys Arg Trp Tyr Gly Trp Lys
                      5
                                          10
      Arg Cys Cys Leu Ser Glu Ser Ser Val Gly Cys Ser Val Ser Trp Ala
                                                          30
                  20
                                      25
30
      Ile Leu Ser Thr Thr Ser Thr Lys Leu Ile Met Ala Val Leu Ser Thr
              35
                                  40
      Ser Ala Thr Met Asn Gly Met Leu Leu Trp Lys Asp Ala Thr Arg Lys
                              55
      Ser Ser Arg Lys Leu Gln Leu Leu Pro His Asp Ser Leu Tyr Leu Phe
35
                          70
                                              75
      Cys Val Pro Gln Gly Leu Lys Val Thr Thr Cys Gly Asp Lys
                      85
      (2) INFORMATION FOR SEQ ID NO:3:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
45
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007804
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
50
      Thr Ile Gly Arg Lys Glu Lys Thr Lys Met Ser Leu Val Trp Leu Glu
                                          10
      Ala Met Leu Pro Leu Gly Ile Ile Gly Gly Met Leu Cys Ile Met Gly
                                      25
      Asn Ser Gln Tyr Tyr Ile His Lys Ala Tyr His Gly Arg Pro Lys His
55
                                  40
      Ile Gly His Asp Glu Trp Asp Val Ala Met Glu Arg Arg Asp Lys
                              55
      Val Val Glu Lys Ala Ala Pro Ser Ser
                          70
60
      (2) INFORMATION FOR SEQ ID NO:4:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 65 amino acids

WO 00/40695

Calculate the second second (C) STRANDEDNESS:

PCT/US00/00466

603

```
(D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 1007805
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
10
      Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile Ile Gly
                     5
                                          10
      Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His Lys Ala
                  20
                                      25
      Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp Val Ala
15
                                  40
      Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Ala Pro Ser
                              55
      Ser
      65
20
      (2) INFORMATION FOR SEO ID NO:5:
           i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 700 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..700
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1008556
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
      aaaagaaaac tgaagaaccc caaaaatccc hagtgaaaga aaacgttatc cacttcgaga
                                                                              60
      gagtgacacg tcacgaagca aatccaaatt Jaaaactttt ccataaaaat caaatcactt
                                                                             120
      cttcttatta cttccccaaa tcgaaatcag tcacttatta aaaccctaaa tcatgacgat
                                                                             180
35
      cgcaccggca ttgcagacga cgttcgtgtc atcaaccaac ttcctgaaac attcttcttc
                                                                             240
      gtggggatca tcatcaccaa acaatgtgat tcttcccaaa aacaagagat cttctacctc
                                                                             300
      cgtagttgtc gccgccgtcg gtgatgtctc ttctgacgga acaatctact taatcggcgg
                                                                             360
      agccatcgcc gttgcactcg tcggaactgc atttccgatc ctcttcaaac gcaaagacac
                                                                             420
      gtgtccggaa tgtgatggag caggatttgt gaggaaagga ggagtgactc tgagagccaa
                                                                             480
40
      cgccgcacgg aaggatcttc ctcagatcgt ttgtgctaat tgcaatggac tcggaaagct
                                                                             540
                                                                             600
      taaccagatt gataaatcat aagtottott catatatatg taaaaagoto catttotoat
                                                                             660
      tcctcttctt atctacaata tacctctgta agagtagata gcttcgaatg atcaatggtt
      gcttggagat ggatggatat atatacaa ttgctttttt
      (2) INFORMATION FOR SEQ ID NO:6:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 129 amino acids
                 (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..129
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1008557
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
      Met Thr Ile Ala Pro Ala Leu Gln Thr Thr Phe Val Ser Ser Thr Asn
                                           10
      Phe Leu Lys His Ser Ser Ser Trp Gly Ser Ser Ser Pro Asn Asn Val
```

25

Ile Leu Pro Lys Asn Lys Arg Ser Ser Thr Ser Val Val Val Ala Ala 35 40 45 Val Gly Asp Val Ser Ser Asp Gly Thr Ile Tyr Leu Ile Gly Gly Ala

20

PCT/US00/00466 WO 00/40695 604 5.5 60 Ile Ala Val Ala Leu Val Gly Thr Ala Phe Pro Ile Leu Phe Lys Arg 75 70 Lys Asp Thr Cys Pro Glu Cys Asp Gly Ala Gly Phe Val Arg Lys Gly 5 85 90 Gly Val Thr Leu Arg Ala Asn Ala Ala Arg Lys Asp Leu Pro Gln Ile 100 105 110 Val Cys Ala Asn Cys Asn Gly Leu Gly Lys Leu Asn Gln Ile Asp Lys 120 10 Ser (2) INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 665 base pairs 15 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 2.0 (A) NAME/KEY: -(B) LOCATION: 1..665 (D) OTHER INFORMATION: / Ceres Seq. ID 1008628 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: atgttactct tctcagatct cttgttcgtt ttgatgtaca atggcgtcct tggtagcagc 25 tectatetet tteteaggtg acteteatgt caaagcacae egaaacttea atgegatteg 120 caagagetet acattgactg ttcaaacaaa atcaaaccge agtcacaaac teteggttte 180 tgcaggttac cgtgggggaa gtaagggtgg tggaagtagt gattttgtta ccggttttct 240 totaggaagt gotgtttcg gaactotggo ttatatottt gotccacaga tocgaagato 300 aqtqctqaqc gaqaatqaat atqqtttcaa qaaaccqqaq caqccqatqt actatqacqa 360 30 aggectagag gagagaagag agatattgaa tgagaaaatc ggccaactca attccgccat 420 tgacaaggtt tcgtcgcgtc tgaaaggagg tcggagcggt agcagcaaga acacttcttc 480 gccgtctgtc ccagttgaaa ccgacgcaga agcagaagct actgcatgat tgaatgtaat 540 cctctgctct attttaccaa ttcaaaactg ccttccattg gttctgtggt ttttttttt 600 tttttgttgg aaccattagg ggcttttctg acttttagat attgaaagaa aaagacaatc 660 35 gtcgc (2) INFORMATION FOR SEO ID NO:8: (i) SEOUENCE CHARACTERISTICS: (A) LENGTH: 162 amino acids (B) TYPE: amino acid 40 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 45 (B) LOCATION: 1..162 (D) OTHER INFORMATION: / Ceres Seq. ID 1008629 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8: Met Ala Ser Leu Val Ala Ala Pro Ile Ser Phe Ser Gly Asp Ser His 5 10 50 Val Lys Ala His Arg Asn Phe Asn Ala Ile Arg Lys Ser Ser Thr Leu 25 Thr Val Gln Thr Lys Ser Asn Arg Ser His Lys Leu Ser Val Ser Ala 40

Thr Val Gln Thr Lys Ser Asn Arg Ser His Lys Leu Ser Val Ser Ala
35
Gly Tyr Arg Gly Gly Ser Lys Gly Gly Gly Ser Ser Asp Phe Val Thr
50
Gly Phe Leu Leu Gly Ser Ala Val Phe Gly Thr Leu Ala Tyr Ile Phe
65
Ala Pro Gln Ile Arg Arg Ser Val Leu Ser Glu Asn Glu Tyr Gly Phe
85
60
Lys Lys Pro Glu Gln Pro Met Tyr Tyr Asp Glu Gly Leu Glu Glu Arg
100
Arg Glu Ile Leu Asn Glu Lys Ile Gly Gln Leu Asn Ser Ala Ile Asp

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605 120 115 Lys Val Ser Ser Arg Leu Lys Gly Gly Arg Ser Gly Ser Ser Lys Asn 135 140 Thr Ser Ser Pro Ser Val Pro Val Glu Thr Asp Ala Glu Ala Glu Ala 5 145 150 155 Thr Ala (2) INFORMATION FOR SEQ ID NO:9: (i) SEQUENCE CHARACTERISTICS: 10 (A) LENGTH: 1101 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) 15 (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..1101 (D) OTHER INFORMATION: / Ceres Seq. ID 1009376 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9: 20 ottotttoot officettott totgfffftot ggtogotogf tacccccaag acctogfoga 60 ggtttctttc tctctgaaaa agtgcttttg ctgcgtttcc atgctctcgg aaggaacttg 120 getgaaregt tggaggatta tegatetgge gtegttggtg catetetage tgegeeaatt 180 categgatee ggtttgattt etecettteg egaatetage etteegtgat ettageeatg 240 ttctatggca cagctgtatg ggaccettgg cttategttg gecagattat ttgcetecaa 300 25 tgctcttact atctcactct tggactcttc accatggtct ttcttggcct tcgtgttcct 360 egecttagte tigictacti citegatiae getactetea etactteeae eticaeeggt 420 tggtctgtta ttgcctcatt cctcttctct tcactcgctg gggctgtgta catgatattt 480 ttggtggagc gagcacggaa atgcttagat ttctctgcaa ctctctacat catacatctc 540 600 ttettttgea teatgtatgg aggatggeet teetetatgg egtggtgggt tgttaatgga 30 660 acceggacteg etgttatgge tttgetaget gagtaettgt geattaaacg egaacagega gagateceta tggategett ceaeteaagg gtttgaagat tggtgaaaga acaagtgatg 720 aagattottg tgggttagca gogogactog tagagcotag tagagcotag tactottaac 780 aattgtgatg tacagagatg aaatcagagg agtagaatgg gcgaaaggaa gttcagctag 840 ccaagcaaca atgttggaca ctggtgcacc aactctgact gcaattttta gtgacttcaa 900 35 atactaagac tttggaatac gtttcaattt gttcttttgt atgaaaaaaa cattagactc 960 gaataggtta ttgatttcat gttgcttctt ttttttgttt atactggatt ttctcttcta 1020 tectetaett ggtgeagett tgtgtgtttt aaaaccagtg teteatteat cataetettt 1080 taccaccaat tgaatccttc c (2) INFORMATION FOR SEQ ID NO:10: 40 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 152 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 45 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..152 (D) OTHER INFORMATION: / Ceres Seq. ID 1009377 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: Met Phe Tyr Gly Thr Ala Val Trp Asp Pro Trp Leu Ile Val Gly Gln 5 10 Ile Ile Cys Leu Gln Cys Ser Tyr Tyr Leu Thr Leu Gly Leu Phe Thr 20 25 30 55 Met Val Phe Leu Gly Leu Arg Val Pro Arg Leu Ser Leu Val Tyr Phe 40 Phe Asp Tyr Ala Thr Leu Thr Thr Ser Thr Phe Thr Gly Trp Ser Val 55 Ile Ala Ser Phe Leu Phe Ser Ser Leu Ala Gly Ala Val Tyr Met Ile

75

90

Phe Leu Val Glu Arg Ala Arg Lys Cys Leu Asp Phe Ser Ala Thr Leu

70

85

606 Tyr Ile Ile His Leu Phe Phe Cys Ile Met Tyr Gly Gly Trp Pro Ser 100 105 110 Ser Met Ala Trp Trp Val Val Asn Gly Thr Gly Leu Ala Val Met Ala 115 120 125 Leu Leu Ala Glu Tyr Leu Cys Ile Lys Arg Glu Gln Arg Glu Ile Pro 130 135 140 Met Asp Arg Phe His Ser Arg Val 150 (2) INFORMATION FOR SEQ ID NO:11: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 120 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..120 (D) OTHER INFORMATION: / Ceres Seq. ID 1009378 20 xi) SEQUENCE DESCRIPTION: SEQ ID NO:11: Met Tal Phe Leu Gly Leu Arg Val Pro Arg Leu Ser Leu Val Tyr Phe 10 Phe Asp Tyr Ala Thr Leu Thr Thr Ser Thr Phe Thr Gly Trp Ser Val 20 25 25 Ile Ala Ser Phe Leu Phe Ser Ser Leu Ala Gly Ala Val Tyr Met Ile 40 Phe Leu Val Glu Arg Ala Arg Lys Cys Leu Asp Phe Ser Ala Thr Leu 55 Tyr Ile Ile His Leu Phe Phe Cys Ile Met Tyr Gly Gly Trp Pro Ser 30 70 75 Ser Met Ala Trp Trp Val Val Asn Gly Thr Gly Leu Ala Val Met Ala 85 90 Leu Leu Ala Glu Tyr Leu Cys Ile Lys Arg Glu Gln Arg Glu Ile Pro 100 105 35 Met Asp Arg Phe His Ser Arg Val 115 (2) INFORMATION FOR SEQ ID NO:12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 76 amino acids 40 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 45 (A) NAME/KEY: peptide (B) LOCATION: 1..76 (D) OTHER INFORMATION: / Ceres Seq. ID 1009379 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12: Met Ala Gln Leu Tyr Gly Thr Leu Gly Leu Ser Leu Ala Arg Leu Phe 50 10 Ala Ser Asn Ala Leu Thr Ile Ser Leu Leu Asp Ser Ser Pro Trp Ser 20 25 30 Phe Leu Ala Phe Val Phe Leu Ala Leu Val Leu Ser Thr Ser Ser Ile 35 40 45 55 Thr Leu Leu Ser Leu Leu Pro Pro Ser Pro Val Gly Leu Leu Pro 50 55 60 His Ser Ser Ser Leu His Ser Leu Gly Leu Cys Thr 70 (2) INFORMATION FOR SEQ ID NO:13: 60 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 398 base pairs (B) TYPE: nucleic acid

607

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(C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
 5
                (A) NAME/KEY: -
                (B) LOCATION: 1..398
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011128
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
      aaatcaataq ettatetagt tettgaetqt teaacgatea atggeaceat eegetgeaat
                                                                              60
10
      gctcatactc tcacatcctc tagttagcca caaagctaag aatcaatctc tgtcatcgcc
                                                                             120
      gtcgtctgtt aagtcgacac gtgtcttcgg ttttctttgg ccttggaagg cattagacaa
                                                                             180
      tgaggatcat tccgcqqttq ttctaqqccq qctctttggc qatccggcta ctatcgagaa
                                                                             240
     gcgcttccaa gaagctcttg aacaaagctg ttggtaattt gtacaatgtt tggttgggtt
                                                                             300
      tattgattgt ttttacgtta aaatcgcttt tataaattgg aaattgaagt actgtaaaat
                                                                             360
15
      gtaaaaattg actatatata attaaaggta cattatgc
      (2) INFORMATION FOR SEQ ID NO:14:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 78 amino acids
                (B) TYPE: amino acid
20
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
25
                (B) LOCATION: 1..78
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011129
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
      Met Ala Pro Ser Ala Ala Met Leu Ile Leu Ser His Pro Leu Val Ser
                      5
                                           10
                                                               15
30
      His Lys Ala Lys Asn Gln Ser Leu Ser Ser Pro Ser Ser Val Lys Ser
                  2.0
                                       25
      Thr Arg Val Phe Gly Phe Leu Trp Pro Trp Lys Ala Leu Asp Asn Glu
                                  4.0
                                                       4.5
      Asp His Ser Ala Val Val Leu Gly Arg Leu Phe Gly Asp Pro Ala Thr
35
                              55
      Ile Glu Lys Arg Phe Gln Glu Ala Leu Glu Gln Ser Cys Trp
                          70
      (2) INFORMATION FOR SEQ ID NO:15:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 72 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
45
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..72
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011130
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:
50
      Met Leu Ile Leu Ser His Pro Leu Val Ser His Lys Ala Lys Asn Gln
                                           10
      Ser Leu Ser Ser Pro Ser Ser Val Lys Ser Thr Arg Val Phe Gly Phe
                                       25
      Leu Trp Pro Trp Lys Ala Leu Asp Asn Glu Asp His Ser Ala Val Val
55
                                   40
      Leu Gly Arg Leu Phe Gly Asp Pro Ala Thr Ile Glu Lys Arg Phe Gln
                               55
      Glu Ala Leu Glu Gln Ser Cys Trp
                          70
60
      (2) INFORMATION FOR SEQ ID NO:16:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 549 base pairs

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608
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
 5
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..549
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011718
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:
10
      agggatgtag taggccaata ggcaaatcag agaatcacaa atggtatctg gtcaagaaga
                                                                              60
      ttcctgtttg tctgcagatg catgtgtgta atctagggta tatgtttttg tccatttggt
      ttcataaggc aataaagatc cagctattta ctacttgtga agaaaaaagt gagggaaaac
      agagtcaagt ccgattactt ttctgggaaa gctgaggatt ttgtgttctt agagagcaga
                                                                             240
      tgaagaggtg actgttgttt ctggtgagat agaagttggc atctaaattt gctttctctg
                                                                             300
15
     gaccagtgaa atcaattgca gtctcctata tattgtagta ggcgatgtat cagtggtgaa
                                                                             360
     qcgaggatga catctgtggg tacttgttct taatccttca ctctgatatc tatgctttag
                                                                             420
      aagcgtttca agttcatgaa gctgatttga tgttgagttt ttaacaacaa gaatcaattc
                                                                             480
      actocaaaaa tatgtaaact ctgccatgcc ttgagctgct ttttggatta tccactgttt
                                                                             540
      tattttata
20
      (2) INFORMATION FOR SEQ ID NO:17:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 70 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
25
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..70
30
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011719
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:
      Gly Cys Ser Arg Pro Ile Gly Lys S€r Glu Asn His Lys Trp Tyr Leu
                      5
                                           10
      Val Lys Lys Ile Pro Val Cys Leu Gln Met His Val Cys Asn Leu Gly
35
                  20
                                      25
      Tyr Met Phe Leu Ser Ile Trp Phe His Lys Ala Ile Lys Ile Gln Leu
                                  40
                                                       45
      Phe Thr Thr Cys Glu Glu Lys Ser Glu Gly Lys Gln Ser Gln Val Arg
                              55
40
      Leu Leu Phe Trp Glu Ser
                          70
      (2) INFORMATION FOR SEQ ID NO:18:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 45 amino acids
45
                 (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
50
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..45
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1011720
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:
      Met His Val Cys Asn Leu Gly Tyr Met Phe Leu Ser Ile Trp Phe His
55
                                           10
      Lys Ala Ile Lys Ile Gln Leu Phe Thr Thr Cys Glu Glu Lys Ser Glu
                                      25
      Gly Lys Gln Ser Gln Val Arg Leu Leu Phe Trp Glu Ser
                                  40
60
      (2) INFORMATION FOR SEQ ID NO:19:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 37 amino acids

TC1/0500/00400

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609
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..37
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011721
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:
10
     Met Phe Leu Ser Ile Trp Phe His Lys Ala Ile Lys Ile Gln Leu Phe
                                         10
     Thr Thr Cys Glu Glu Lys Ser Glu Gly Lys Gln Ser Gln Val Arg Leu
                 20
                                      25
     Leu Phe Trp Glu Ser
15
              35
      (2) INFORMATION FOR SEQ ID NO:20:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 417 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..417
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011735
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:
      aattoottot tittoacato tacccaaaat tocaaaacac categatitt tigotototi
                                                                            60
      gaaatgctct ttttatgtcc aattcttctc ttccaactaa accgaaccgg aaaaccaggt
                                                                            120
                                                                          180
30
      teggagateg gtgtttaetg atggecaaac ageaacgaac eegaetttae atacteegaa
      gatgtgtctc catgctgctt tgctggcacg accactctat ttctgattag tttgcagtca
                                                                            240
      tgaaccggac tgaccaagaa gtagcaatca gaagaatgtg atgtcattta gagttttttc
                                                                            300
      ttgttttctt ttagaactct cttcagggaa tcttttgtaa ttgaagaaga ctctaaggat
                                                                           360
      tgggcctttt gggcctttgt acatattgtg taaataacat atgaactttt tttaccc
35
      (2) INFORMATION FOR SEQ ID NO:21:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 51 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
40
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..51
45
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011736
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:
      Met Ser Asn Ser Ser Leu Pro Thr Lys Pro Asn Arg Lys Thr Arg Phe
                     5
                                         10
      Gly Asp Arg Cys Leu Leu Met Ala Lys Gln Gln Arg Thr Arg Leu Tyr
50
                 20 25
                                                          30
      Ile Leu Arg Arg Cys Val Ser Met Leu Leu Cys Trp His Asp His Ser
             3.5
                                 40
      Ile Ser Asp
55
      (2) INFORMATION FOR SEQ ID NO:22:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 46 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
60
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
```

(ix) FEATURE:

610

```
(A) NAME/KEY: peptide
               (B) LOCATION: 1..46
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011737
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:
 5
     Asn Ser Phe Phe Phe Thr Ser Thr Gln Asn Ser Lys Thr Pro Ser Ile
                                         10
     Phe Cys Ser Leu Glu Met Leu Phe Leu Cys Pro Ile Leu Leu Phe Gln
                                     25
     Leu Asn Arg Thr Gly Lys Pro Gly Ser Glu Ile Gly Val Tyr
10
                                 40
      (2) INFORMATION FOR SEQ ID NO:23:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 32 amino acids
               (B) TYPE: amino acid
1.5
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
20
               (B) LOCATION: 1..32
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011738
          'xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:
      Ile Pro Ser Phe Ser His Leu Pro Lys Ile Pro Lys His His Arg Phe
                                         10
25
     Phe Ala Leu Leu Lys Cys Ser Phe Tyr Val Gln Phe Phe Ser Ser Asn
                 20
                                     25
      (2) INFORMATION FOR SEQ ID NO:24:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 712 base pairs
30
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
35
                (A) NAME/KEY: -
                (B) LOCATION: 1..712
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011755
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:
      actcaaatag atgaagaaat ggcgagctta agcacgagct tgctcggaat aagacgaggg
                                                                           60
40
      tcaagtaaca agaacaaggc agaagataag accaaaaacc agtgttttgc aagaagcatc
                                                                           120
      caaagcatag acaatcacct ggtgtttgtt ctctttgtct taacgagagg ctctctcttt
                                                                           180
      ttatcaaagc agcttcttca cgtagaccaa gatctcgtca aatcttgtct acttcttcct
                                                                           240
                                                                           300
      caactacttc ctctcttca tctgacggtt cttcttctgt ctcttcttgt ccttctccta
                                                                          360
      ttgttgatcg ccgacqatac ttgttaatgt ccggaggcag cggtagagga gagaaagtga
45
                                                                          420
      tttcgtggat gacgaaaagc cgatccgtag cttataaagt ggatgatgag aaaagaagga
      agaagaagac aaagacgaat agtggtttct tttttggttt ggtaatgggc acaaagaaga
                                                                          480
                                                                          540
      tettgattet tteattttag attgattget acatatagat gaetagatea gtttggtggg
                                                                          600
      tacgtgcaag tgagagagat tetttatacg tattettgtg attgatteee taagettgta
                                                                          660
50
      qtttagtgtg gtaacataaa atgtgataat ccagatttaa tatgcgttga ac
      (2) INFORMATION FOR SEQ ID NO:25:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 108 amino acids
                (B) TYPE: amino acid
55
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
60
                (B) LOCATION: 1..108
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011756
```

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

611 Thr Gln Ile Asp Glu Glu Met Ala Ser Leu Ser Thr Ser Leu Leu Gly 10 Ile Arg Arg Gly Ser Ser Asn Lys Asn Lys Ala Glu Asp Lys Thr Lys 25 20 Asn Gln Cys Phe Ala Arg Ser Ile Gln Ser Ile Asp Asn His Leu Val 40 35 Phe Val Leu Phe Val Leu Thr Arg Gly Ser Leu Phe Leu Ser Lys Gln 55 Leu Leu His Val Asp Gln Asp Leu Val Lys Ser Cys Leu Leu Pro 10 75 Gln Leu Leu Pro Leu Phe His Leu Thr Val Leu Leu Ser Leu Leu 90 Val Leu Leu Leu Leu Ile Ala Asp Asp Thr Cys 100 (2) INFORMATION FOR SEQ ID NO:26: 15 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 102 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..102 25 (D) OTHER INFORMATION: / Ceres Seq. ID 1011757 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26: Met Ala Ser Leu Ser Thr Ser Leu Leu Gly Ile Arg Arg Gly Ser Ser 1.0 Asn Lys Asn Lys Ala Glu Asp Lys Thr Lys Asn Gln Cys Phe Ala Arg 30 20 25 Ser Ile Gln Ser Ile Asp Asn His Leu Val Phe Val Leu Phe Val Leu Thr Arg Gly Ser Leu Phe Leu Ser Lys Gln Leu Leu His Val Asp Gln 35 Asp Leu Val Lys Ser Cys Leu Leu Pro Gln Leu Pro Leu Phe 70 His Leu Thr Val Leu Leu Ser Leu Leu Val Leu Leu Leu Leu 85 Ile Ala Asp Asp Thr Cys 40 100 (2) INFORMATION FOR SEQ ID NO:27: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 422 base pairs (B) TYPE: nucleic acid 45 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -50 (B) LOCATION: 1..422 (D) OTHER INFORMATION: / Ceres Seq. ID 1011832 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27: aataacaata tcaaaaqata aaaacaqagt ttqcttttat aagatcaaag aaggtaaaaa 60 120 aaaaaagtgg caaaatggag tgcgatcgta ttgataatga tggtgatgat tgttgcggtc 55 acaatagaag cacaagaaga aagtggttgg actatttgtt ttcgccaatg ttctcagcct 180 240 tgtcgcagtg acgacggtag ttgctatgaa aactgtaaaa tcgaatgcgg tggtcccaag 300 cctcccttat ctcqtctacq aagctcacat qcatqaqatq qcatcatcaa tqqaaqttcq 360 cqqaaqqata qgatqataaa aaqaqatttq taqcqqattt ttqatcatct ctactqtttt taacttcacq ttttatatta taaqaqtttc attaaaaaqa tcaataaaac qaqaaaatqt 420 60 (2) INFORMATION FOR SEQ ID NO:28: (i) SEQUENCE CHARACTERISTICS:

612

```
(A) LENGTH: 59 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..59
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011833
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:
     Met Met Val Met Ile Val Ala Val Thr Ile Glu Ala Gln Glu Glu Ser
                     5
                                          10
     Gly Trp Thr Ile Cys Phe Arg Gln Cys Ser Gln Pro Cys Arg Ser Asp
                                                          30
                 20
                                      25
15
     Asp Gly Ser Cys Tyr Glu Asn Cys Lys Ile Glu Cys Gly Gly Pro Lys
                                 40
      Pro Pro Leu Ser Arg Leu Arg Ser Ser His Ala
                              55
      (2) INFORMATION FOR SEQ ID NO:29:
20
            i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 58 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
25
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..58
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011834
30
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:
      Met Val Met Ile Val Ala Val Thr Ile Glu Ala Gln Glu Glu Ser Gly
                                          10
      Trp Thr Ile Cys Phe Arg Gln Cys Ser Gln Pro Cys Arg Ser Asp Asp
                                  25
                 20
                                                          30
35
      Gly Ser Cys Tyr Glu Asn Cys Lys Ile Glu Cys Gly Gly Pro Lys Pro
             35
                                40
      Pro Leu Ser Arg Leu Arg Ser Ser His Ala
                             55
      (2) INFORMATION FOR SEQ ID NO:30:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 56 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..56
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011835
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:
      Met Ile Val Ala Val Thr Ile Glu Ala Gln Glu Glu Ser Gly Trp Thr
                     5
                                          10
      Ile Cys Phe Arg Gln Cys Ser Gln Pro Cys Arg Ser Asp Asp Gly Ser
                                     25
55
      Cys Tyr Glu Asn Cys Lys Ile Glu Cys Gly Gly Pro Lys Pro Pro Leu
                                  40
      Ser Arg Leu Arg Ser Ser His Ala
       (2) INFORMATION FOR SEQ ID NO:31:
60
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 713 base pairs
```

(B) TYPE: nucleic acid

613

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(C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
 5
               (A) NAME/KEY: -
               (B) LOCATION: 1..713
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011907
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:
     aattotggtt tttettgcaa tgettettee atttetetee teetetete teeteegtet
                                                                            60
10
     ctcttctccq atcatqtccc ctqatcccaa atccaatacc ttcttcttct ctttttccaa
                                                                           120
     ttgattaccg gagaatttca gatcgcgatc tccgatttat taataattac agaaaaaaat
                                                                            180
     aattaaaccc tagagagaga gagagatata tatatacatg gagagttcat taggtttcat
                                                                            240
     ggcggttttc gccgtctcag gaagcgttgt gttcttagcg agtcaatttc acaagcgtct
                                                                            300
     tototogat tacatggaca agttogaatt cgaaatccga gcgcagaaaa aaatggtgat
                                                                            360
15
     qaaqaaqaag gtgagattcg cggcggatgt ggtggagccg tcggggaata acaaaqagta
                                                                            420
     togooggaga cartottoca aggotaaato gaattogaag atggoggoaa ctatttgact
                                                                            480
     ttaaggtttt ttgtacaaaa tttaatgtgt aatttattca tttgggtttt tgtgatttga
                                                                            540
     aattcgtaaa tttaatttcg gaatctgaat ctgggctaaa cttttcaagt cttccccagg
                                                                            600
      gtcaattctt tttcttcttc tttttttct tgaggggttt ttttttgttg ttgtgacctg
                                                                            660
20
      taaataaatg tgttgtttgt gtatatacaa aatgtgattg ctaattacct ttt
      (2) INFORMATION FOR SEQ ID NO:32:
           (1) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 86 amino acids
                (B) TYPE: amino acid
25
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
30
                (B) LOCATION: 1..86
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011908
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:
      Met Glu Ser Ser Leu Gly Phe Met Ala Val Phe Ala Val Ser Gly Ser
                                         10
                      5
35
      Val Val Phe Leu Ala Ser Gln Phe His Lys Arg Leu Leu Ser Asp Tyr
                  20
                                     25
      Met Asp Lys Phe Glu Phe Glu Ile Arg Ala Gln Lys Lys Met Val Met
                                  40
      Lys Lys Val Arg Phe Ala Ala Asp Val Val Glu Pro Ser Gly Asn
40
                          55
                                                 60
      Asn Lys Glu Tyr Arg Arg His Ser Ser Lys Ala Lys Ser Asn Ser
                         70
      Lys Met Ala Ala Thr Ile
                      85
45
      (2) INFORMATION FOR SEQ ID NO:33:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 79 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
50
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..79
55
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011909
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:
      Met Ala Val Phe Ala Val Ser Gly Ser Val Val Phe Leu Ala Ser Gln
                                          10
      Phe His Lys Arg Leu Leu Ser Asp Tyr Met Asp Lys Phe Glu Phe Glu
60
                                  25
      Ile Arg Ala Gln Lys Lys Met Val Met Lys Lys Val Arg Phe Ala
```

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Ala Asp Val Val Glu Pro Ser Gly Asn Asn Lys Glu Tyr Arg Arg
                              55
     His Ser Ser Lys Ala Lys Ser Asn Ser Lys Met Ala Ala Thr Ile
                          70
5
      (2) INFORMATION FOR SEQ ID NO:34:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 59 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
10
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..59
15
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011910
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:
      Phe Trp Phe Phe Leu Gln Cys Phe Phe His Phe Ser Pro Pro Leu Ser
                      5
                                          10
      Ser Ser Val Ser Leu Leu Arg Ser Cys Pro Leu Ile Pro Asn Pro Ile
20
                  20
                                      25
      Pro Ser Ser Ser Leu Phe Fro Ile Asp Tyr Arg Arg Ile Ser Asp Arg
                                                       45
                                  40
      Asp Leu Arg Phe Ile Asn Asn Tyr Arg Lys Lys
                              55
25
      (2) INFORMATION FOR SEQ ID NO:35:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 580 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
30
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..580
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011911
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:
      aattgtttta aaattacaaa ttagtccgtt cttttattcc cgtactcgtt ccttcttctt
                                                                              60
      cttcttcctc tcatcgtcat tttctcgatt ctcactcttc cggtcaccga ctaattctga
                                                                             120.
      ataaggttta tcaaaagaat aagaataagt ggataaaaag ctagctttga aagagttatt
40
      gcagagaaaa aaaatgggat cgagagggat tatcaacgat aagtggtcaa tgaggattct
                                                                             240
      atggggttgt gctatcggaa gtgctattgg tttatacatg gttgctgtag agagacaaac
                                                                             300
      tcagaacagg gctcgtgcta tggctgagag tttgagagct gctgaatcac aaggtgatgg
                                                                             360
      tgataatgtc taatatctac caagtagtgc tcagttgaat actctcagtt gagttttttt
                                                                             420
      ttttggtgtt tgtttttgtt ataatgactt cttctgccaa gatggtgttg atgtagtttc
                                                                             480
45
      ttttttgcaa ataatcgtaa taaqgtttcg aaacttggag agttgaagtt gctgaacata
                                                                             540
      cgatttgtgt tatcgcaaaa aaagttattt cttatgcctg
      (2) INFORMATION FOR SEQ ID NO:36:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 59 amino acids
50
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
55
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..59
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011912
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:
      Met Gly Ser Arg Gly Ile Ile Asn Asp Lys Trp Ser Met Arg Ile Leu
60
                                           10
      Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr Met Val Ala Val
                   20
                                       25
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Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala Glu Ser Leu Arg
                                  40
             35
     Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
                              55
5
      (2) INFORMATION FOR SEQ ID NO:37:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 47 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
10
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..47
15
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011913
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:
      Met Arg Ile Leu Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr
                                          10
      Met Val Ala Val Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala
20
                  20
                                      25
      Glu Ser Leu Arg Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
                                  40
      (2) INFORMATION FOR SEQ ID NO:38:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 40 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..40
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011914
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:
35
      Ile Val Leu Lys Leu Gln Ile Ser Pro Phe Phe Tyr Ser Arg Thr Arg
                                           10
      Ser Phe Phe Phe Phe Leu Ser Ser Ser Phe Ser Arg Phe Ser Leu
                  20
                                      25
      Phe Arg Ser Pro Thr Asn Ser Glu
40
              35
      (2) INFORMATION FOR SEQ ID NO:39:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 415 base pairs
                (B) TYPE: nucleic acid
45
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
50
                (B) LOCATION: 1..415
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011954
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:
      tcacaaaacc tgattgttct gtttcgaacc gatagattcg aaccttggtt taagttgccg
                                                                              60
      gcttcgagga atctccggtg tggactttca ctcctgtttc ttcttcatga cgcaagctgt
                                                                             120
55
      tttctctaga atacgctttc ttagaaccct ggaaaacgtt acccagcttc ctcaatcctc
                                                                             180
      tagetetage tegaagaaca teetttgaaa caettggate tttaagcace gettggeaga
                                                                             240
                                                                            300
      cacqagataa qqttqattcq atqtcaacaa catttatttq ccaaaqqqat tqaaqcattq
      catecttttt agetteaatg getttetgaa tatgtetetg atetttgtga ttgtgatgat
                                                                            360
      catgcatgte ctategttge gatgtatttt ataaatgtte gtetgtaagt tattt
60
      (2) INFORMATION FOR SEQ ID NO:40:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 42 amino acids
```

```
616
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..42
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011955
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:
10
     Ser Gln Asn Leu Ile Val Leu Phe Arg Thr Asp Arg Phe Glu Pro Trp
                     5
                                          10
      Phe Lys Leu Pro Ala Ser Arg Asn Leu Arg Cys Gly Leu Ser Leu Leu
                  20
                                      .25
      Phe Leu His Asp Ala Ser Cys Phe Leu
15
              35
                                  40
      (2) INFORMATION FOR SEQ ID NO:41:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 33 amino acids
                (B) TYPE: amino acid
20
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
25
                (B) LOCATION: 1..33
                (D) OTHER INFORMATION: / Ceres Seq. ID 1011956
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:
      Met Thr Gln Ala Val Phe Ser Arg Ile Arg Phe Leu Arg Thr Leu Glu
                   5
                                         10
                                                              15
30
      Asn Val Thr Gln Leu Pro Gln Ser Ser Ser Ser Ser Lys Asn Ile
                  20
                                      25
                                                           30
      Leu
      (2) INFORMATION FOR SEQ ID NO:42:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 440 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..440
```

(D) OTHER INFORMATION: / Ceres Seq. ID 1011960

45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:

> gacaqtcaqt cactqtaaca ttttagatct ttcccqaaqa aqaaaacqaa gaaqaqacqa 60 agagagaaat gaggccgatg cagctggata tgttatcgga gatggatgat gcaggttctt 120 cgatggccat ggacgttgat gacctcgaag ccatggagat actcaacgaa ggaggacttg 180 totcagataa caagotogoo gacqooqatt tottcaacaa attogatqat qatttogatq 240 acaccgatat caactaaatc cggttaagtt ttctctttaa gatttcagtt tttttccta 300 aacttgatgt aatttgtgga gaaacctatg tagtaagtgt gtaataagtt ctgggatttt 360 tatgtttgtg gtatgtaaca atgtttttct tcatttggat tactaaaaca tttgatttgt 420 gtgttattgt gctttgtctg

(2) INFORMATION FOR SEO ID NO:43:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 62 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
  - (D) TOPOLOGY: linear
- 60 (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:

50

55

(A) NAME/KEY: peptide

```
(B) LOCATION: 1..62
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011961
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:
     Met Arg Pro Met Gln Leu Asp Met Leu Ser Glu Met Asp Asp Ala Gly
 5
              5
                                     10
     Ser Ser Met Ala Met Asp Val Asp Asp Leu Glu Ala Met Glu Ile Leu
                                     25
     Asn Glu Gly Gly Leu Val Ser Asp Asn Lys Leu Ala Asp Ala Asp Phe
                          40
10
     Phe Asn Lvs Phe Asp Asp Phe Asp Asp Thr Asp Ile Asn
                            55
      (2) INFORMATION FOR SEQ ID NO:44:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 59 amino acids
15
               (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
20
               (A) NAME/KEY: peptide
               (B) LOCATION: 1..59
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011962
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:
     Met Gln Leu Asp Met Leu Ser Glu Met Asp Asp Ala Gly Ser Ser Met
25
          5
                                  10
     Ala Met Asp Val Asp Asp Leu Glu Ala Met Glu Ile Leu Asn Glu Gly
                             25
     Gly Leu Val Ser Asp Asn Lys Leu Ala Asp Ala Asp Phe Phe Asn Lys
                                40
30
      Phe Asp Asp Phe Asp Asp Thr Asp Ile Asn
                     55
      (2) INFORMATION FOR SEQ ID NO:45:
          (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 55 amino acids
35
                (B) TYPE: amino acid
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..55
               (D) OTHER INFORMATION: / Ceres Seq. ID 1011963
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:
      Met Leu Ser Glu Met Asp Asp Ala Gly Ser Ser Met Ala Met Asp Val
45
                                         10
      Asp Asp Leu Glu Ala Met Glu Ile Leu Asn Glu Gly Gly Leu Val Ser
                20
                                    25
      Asp Asn Lys Leu Ala Asp Ala Asp Phe Phe Asn Lys Phe Asp Asp Asp
            35
                          40
50
      Phe Asp Asp Thr Asp Ile Asn
      (2) INFORMATION FOR SEQ ID NO:46:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 193 base pairs
55
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
60
                (A) NAME/KEY: -
                (B) LOCATION: 1..193
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014075
```

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:
     aaaaccctta agctaagcaa atttcaaagc aattcaaaaa ccctagcctc tctctttta
      tctctcacaa caaaatcttt gaaatggcga tctctaaggc ttccattgtt gttctcatga
      tggtgattat ctccgtcgtt gcatcggcac agtctgaggc accagcacca agtcctactt
                                                                             180
 5
     ctggatctag tcg
      (2) INFORMATION FOR SEQ ID NO:47:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 39 amino acids
                (B) TYPE: amino acid
10
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
15
                (B) LOCATION: 1..39
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014076
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:
      Lys Thr Leu Lys Leu Ser Lys Phe Gln Ser Asn Ser Lys Thr Leu Ala
                  5
                                          10
20
      Ser Leu Ser Leu Ser Leu Thr Thr Lys Ser Leu Lys Trp Arg Ser Leu
                                      25
                  20
                                                           30
      Arg Leu Pro Leu Leu Phe Ser
              35
      (2) INFORMATION FOR SEQ ID NO:48:
25
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 36 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..36
                (D) OTHER INFORMATION: / Ceres Seq. ID 1014077
35
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:
      Met Ala Ile Ser Lys Ala Ser Ile Val Val Leu Met Met Val Ile Ile
                                          10
      Ser Val Val Ala Ser Ala Gln Ser Glu Ala Pro Ala Pro Ser Pro Thr
                  20
                                       25
40
      Ser Gly Ser Ser
              35
      (2) INFORMATION FOR SEQ ID NO:49:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 711 base pairs
45
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
50
                (A) NAME/KEY: -
                 (B) LOCATION: 1..711
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1015865
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:
      actetteegg teacegacta attetggtta geteacatte ettatgetea tteatecaca
                                                                              60
55
      gttccctaat taacgctacg aattttgttt ctcctcttat ttcttctttc ctcttgtaga
                                                                             120
      ataaggttta tcaaaagaat aagaataagg tttatctctt aaatctccga taattagcag
                                                                             180
      agttttttca ggattgagtt tgatttgctg ttttggaatc ataaattttg cgttttagtg
                                                                             240
      gataaaaaagc tagcttaaag agttattgca gagaaaaaaa atgggatcga gagggattat
                                                                             300
      caacgataag tggtcaatga ggattctatg gggttgtgct atcggaagtg ctattggttt
                                                                             360
60
                                                                             420
      atacatqqtt qctqtaqaga qacaaactca qaacaqqqct cqtqctatqg ctqaqaqttt
      gagagetget gaateacaag gtgatggtga taatgtetaa tatetaceaa gtagtgetea
                                                                              480
      gttgaatact ctcagttgag tttttttttt tggtgttttgt ttttgttata atgacttctt
                                                                              540
```

```
600
      ctgccaagat ggtgttgatg tagtttcttt tttgcaaata atcgtaataa ggtttcgaaa
      cttggagagt tgaagttgct gaacatacga tttgtgttat cgcaaaaaaa gttatttctt
                                                                             660
      atgcctgtca tgctatgttt gtgaattcga tttttaatgc gtattttcag c
      (2) INFORMATION FOR SEQ ID NO:50:
 5
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 59 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
10
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..59
                (D) OTHER INFORMATION: / Ceres Seq. ID 1015866
15
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:
      Met Gly Ser Arg Gly Ile Ile Asn Asp Lys Trp Ser Met Arg Ile Leu
                                          10
                      5
      Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr Met Val Ala Val
                  20
                                       25
20
      Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala Glu Ser Leu Arg
                                  40
      Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
                              55
          50
      (2) INFORMATION FOR SEQ ID NO:51:
25
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 47 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
30
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..47
                (D) OTHER INFORMATION: / Ceres Seq. ID 1015867
35
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:
      Met Arg Ile Leu Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr
                                           1.0
      Met Val Ala Val Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala
                                       25
      Glu Ser Leu Arg Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
40
              35
                                  40
      (2) INFORMATION FOR SEQ ID NO:52:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 38 amino acids
45
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
50
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..38
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1015868
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:
      Ser Ser Gly His Arg Leu Ile Leu Val Ser Ser His Ser Leu Cys Ser
55
                                           10
      Phe Ile His Ser Ser Leu Ile Asn Ala Thr Asn Phe Val Ser Pro Leu
                                       25
      Ile Ser Ser Phe Leu Leu
               35
60
       (2) INFORMATION FOR SEQ ID NO:53:
            (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 432 base pairs

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620
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
 5
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..432
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021371
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:
10
      gtcattttct cgattctcac tcttccggtc accgactaat tctgagaaaa aaaatgggat
                                                                              60
      cgagagggat tatcaacgat aagtggtcaa tgaggattct atggggttgt gctatcggaa
                                                                             120
     gtgctattgg tttatacatg gttgctgtag agagacaaac tcagaacagg gctcgtgcta
                                                                             180
                                                                             240
      tggctgagag tttgagagct gctgaatcac aaggtgatgg tgataatgtc taatatctac
                                                                             300
      caagtagtgc tcaqttgaat actctcagtt gagttttttt ttttgqtgtt tgtttttgtt
15
      ataatgactt cttctgccaa gatggtgttg atgtagtttc ttttttgcaa ataatcgtaa
                                                                             360
      taaqqtttcq aaacttqqaq aqttqaaqtt qctqaacata cqatttqtqt tatcqcaaaa
                                                                             420
      aaagttattt cc
      (2) INFORMATION FOR SEQ ID NO:54:
           (i) SEQUENCE CHARACTERISTICS:
20
                (A) LENGTH: 76 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
25
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..76
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021372
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:
30
      His Phe Leu Asp Ser His Ser Ser Gly His Arg Leu Ile Leu Arg Lys
                                           10
      Lys Met Gly Ser Arg Gly Ile Ile Asn Asp Lys Trp Ser Met Arg Ile
                                       25
      Leu Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr Met Val Ala
35
                                   40
      Val Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala Glu Ser Leu
                               55
      Arg Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
                          70
40
      (2) INFORMATION FOR SEQ ID NO:55:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 59 amino acids
                (B) TYPE: amino acid(C) STRANDEDNESS:
45
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..59
50
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1021373
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:
      Met Gly Ser Arg Gly Ile Ile Asn Asp Lys Trp Ser Met Arg Ile Leu
                                           10
      Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr Met Val Ala Val
55
                                       25
                  20
                                                            30
      Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala Glu Ser Leu Arg
                                   4 0
      Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
                               55
60
      (2) INFORMATION FOR SEQ ID NO:56:
            (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 47 amino acids

WO 00/40695

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621
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..47
                (D) OTHER INFORMATION: / Ceres Seq. ID 1021374
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:
10
     Met Arg Ile Leu Trp Gly Cys Ala Ile Gly Ser Ala Ile Gly Leu Tyr
                                          1.0
     Met Val Ala Val Glu Arg Gln Thr Gln Asn Arg Ala Arg Ala Met Ala
                  20
                                      25
      Glu Ser Leu Arg Ala Ala Glu Ser Gln Gly Asp Gly Asp Asn Val
15
                                  40
      (2) INFORMATION FOR SEQ ID NO:57:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 286 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..286
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022578
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:
      caccattgga tggttcggct atacaagctg aaacctccca ggaacaggat tcgtggtaaa
                                                                             60
      gcaaagaaac tgaaatcgaa aacaagctct ggattgagtt caaaatcagc gaagaagaac
                                                                             120
30
      ccgtgggtct agaaatccta gaattttcaa tgttggatgg tgagttatat gtcaactctt
                                                                             180
                                                                             240
      aggtetettt attategaga aagttgttge agaegeagte aagtttetgg tgtaggatte
      gtaaaaattt tgttaacttt attcgaattt catgtttatt gcggag
      (2) INFORMATION FOR SEQ ID NO:58:
           (i) SEQUENCE CHARACTERISTICS:
35
                (A) LENGTH: 43 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
40
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..43
                (D) OTHER INFORMATION: / Ceres Seq. ID 1022579
          (xi) SEQUENCE DESCRIPTION: SEO ID NO:58:
45
      His His Trp Met Val Arg Leu Tyr Lys Leu Lys Pro Pro Arg Asn Arg
                                          10
      Ile Arg Gly Lys Ala Lys Lys Leu Lys Ser Lys Thr Ser Ser Gly Leu
                 20
                                      25
      Ser Ser Lys Ser Ala Lys Lys Asn Pro Trp Val
50
              35
      (2) INFORMATION FOR SEQ ID NO:59:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 43 amino acids
                (B) TYPE: amino acid
55
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
60
                 (B) LOCATION: 1..43
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1022580
```

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:

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622
      Met Val Ser Tyr Met Ser Thr Leu Arg Ser Leu Tyr Tyr Arg Glu Ser
                    5
                                             10
      Cys Cys Arg Arg Ser Gln Val Ser Gly Val Gly Phe Val Lys Ile Leu
                 20
                                         25
 5
      Leu Thr Leu Phe Glu Phe His Val Tyr Cys Gly
      (2) INFORMATION FOR SEO ID NO:60:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 40 amino acids
10
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide.
           ix) FEATURE:
15
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..40
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1022581
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:
      Met Val Arg Leu Tyr Lys Leu Lys Pro Pro Arg Asn Arg Ile Arg Gly
20
                                              10
      Lys Ala Lys Lys Leu Lys Ser Lys Thr Ser Ser Gly Leu Ser Ser Lys
                   20
                                          25
      Ser Ala Lys Lys Asn Pro Trp Val
               35
25
      (2) INFORMATION FOR SEO ID NO:61:
            (i) SEQUENCE CHARACTERISTICS:
                  (A) LENGTH: 643 base pairs
                  (B) TYPE: nucleic acid
                  (C) STRANDEDNESS: single
30
                  (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: DNA (genomic)
           (ix) FEATURE:
                  (A) NAME/KEY: -
                  (B) LOCATION: 1..643
35
                  (D) OTHER INFORMATION: / Ceres Seq. ID 1024240
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:
      agactcagtg aggattttct aggcgatttt ctcgagaaaa tcgtttgagg aaaatggaga
                                                                                    60
                                                                                    120
      cttctatgag gtataccagc aattccaagt ctatgaagat tcatgccaaa gagaaggttc
                                                                                    180
      cggtgaactc aaaaacccat ttacagcttc atggagagtt agatactgga actggggctc
40
                                                                                    240
      cgagttactt ctgtgcgatg attagacact tttttcctga ggccttgggg taggattgca
      ttatgataag cgccaaaagc ttcggtgtct tgtacgcgga aaaaaagagt ttcctgtaag
                                                                                   300
                                                                                   360
      agctgataag cgtgtaacct ttaatattaa agggcqgtgt gatattgatc aggacttaaa
      tcagaagaac cccaaaggag cagcagaatt tgcctggaac ataatggatt tcaaggaaga tcaggatgta cggatcaaag ttggctacga aatgtttgat aaggtccctt atatgcagat tagagaaaac aattggactc tcaacgcgaa catgaaggga aaatggaact tgcggtatga cctgtaactg cattttttc aatcatcatc tgagaaatgt attgatacca ctgctgatga
                                                                                   420
                                                                                   480
45
                                                                                   540
                                                                                   600
      acacatttta attctaccaa ttaatcaaat tcagagatct tcc
       (2) INFORMATION FOR SEQ ID NO:62:
            (i) SEQUENCE CHARACTERISTICS:
50
                  (A) LENGTH: 59 amino acids
                  (B) TYPE: amino acid
                  (C) STRANDEDNESS:
                  (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
55
           (ix) FEATURE:
                  (A) NAME/KEY: peptide
                  (B) LOCATION: 1..59
                  (D) OTHER INFORMATION: / Ceres Seq. ID 1024241
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:
60
      Met Glu Thr Ser Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile
                                              10
       His Ala Lys Glu Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu
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623
                                    25
                 20
                                                         30
     His Gly Glu Leu Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala
                          40
     Met Ile Arg His Phe Phe Pro Glu Ala Leu Gly
 5
      (2) INFORMATION FOR SEQ ID NO:63:
           (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 55 amino acids
               (B) TYPE: amino acid
10
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
               (A) NAME/KEY: peptide
15
               (B) LOCATION: 1..55
               (D) OTHER INFORMATION: / Ceres Seq. ID 1024242
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:
     Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile His Ala Lys Glu
                                         10
20
      Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu His Gly Glu Leu
                                     25
     Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala Met Ile Arg His
                         40
     Phe Phe Pro Glu Ala Leu Gly
25
                         55
       50
      (2) INFORMATION FOR SEQ ID NO:64:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 47 amino acids
                (B) TYPE: amino acid
30
               (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
35
                (B) LOCATION: 1..47
               (D) OTHER INFORMATION: / Ceres Seq. ID 1024243
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:
      Met Asp Phe Lys Glu Asp Gln Asp Val Arg Ile Lys Val Gly Tyr Glu
                                        10
40
      Met Phe Asp Lys Val Pro Tyr Met Gln Ile Arg Glu Asn Asn Trp Thr
                20 . 25
      Leu Asn Ala Asn Met Lys Gly Lys Trp Asn Leu Arg Tyr Asp Leu
      (2) INFORMATION FOR SEQ ID NO:65:
45
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 729 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
50
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..729
                (D) OTHER INFORMATION: / Ceres Seq. ID 1026562
55
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:
      gttaatcata taaatttgac cctaaaatct atctcattct ctcatcaatc ttcttccttt
      aaatccctaa atcttcacgt tacttttcta tctacatttt cactttcaat ggctttgatt
                                                                           120
      atcacctgct cogetetace caegateega geatetteeg gateeggate cettaacceg
                                                                           180
      gatcaaaacc gtaagaaatc tgctqcttgg tgggctcctc tcttcggttt accctccgat
                                                                           240
60
      ccagattacc tcaacatcga aagctcatgc tccaccgtga atccggataa aaccgatatt
                                                                           300
      tccgggtcgg gtcaaaagtt tcgtcgcggt tgttttacgg aggagaaagc taagcagttg
                                                                           360
      aggaggaaaa ccgcagaagc ttccacgttc catgacgtaa tgtatcactc cgccattgct
                                                                           420
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624

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totogacttg cgtotgatat caccggccga gtcgaggatt gaaacggatt cgggtcagat
                                                                            480
     ctggagacga atcaattaat gaataatgaa ttaatctttt ttttcttaat tcagtgttct
                                                                            540
                                                                            600
     tgtaggatgc agaccatctt ctatcggtgc tcctttttca tttgaaccgt tggcttgtgt
     ttggtctttg tgtgttgtaa actctggttc cctttctgtt tcttctgtaa tcaaccgttg
                                                                            660
 5
                                                                            720
     gatttccact tgttattttt agacatgtgt tccctagtta gctcttttat cttatctcaa
     (2) INFORMATION FOR SEQ ID NO:66:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 153 amino acids
10
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
15
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..153
                (D) OTHER INFORMATION: / Ceres Seq. ID 1026563
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:
     Val Asn His Ile Asn Leu Thr Leu Lys Ser Ile Ser Phe Ser His Gln
20
                      5
                                          10
     Ser Ser Ser Phe Lys Ser Leu Asn Leu His Val Thr Fhe Leu Ser Thr
                  20
                                      25
                                                          30
      Phe Ser Leu Ser Met Ala Leu Ile Ile Thr Cys Ser Ala Leu Pro Thr
25
      Ile Arg Ala Ser Ser Gly Ser Gly Ser Leu Asn Pro Asp Gln Asn Arg
                              55
     Lys Lys Ser Ala Ala Trp Trp Ala Pro Leu Phe Gly Leu Prc Ser Asp
                          7.0
                                              75
      Pro Asp Tyr Leu Asn Ile Glu Ser Ser Cys Ser Thr Val Asn Pro Asp
30
                                          90
     Lys Thr Asp Ile Ser Gly Ser Gly Gln Lys Phe Arg Arg Gly Cys Phe
                  100
                                      105
                                                          110
     Thr Glu Glu Lys Ala Lys Gln Leu Frg Arg Lys Thr Ala Glu Ala Ser
                                 120
             115
                                                     125
35
     Thr Phe His Asp Val Met Tyr His Ser Ala Ile Ala Ser Arg Leu Ala
                             135
                                                  140
     Ser Asp Ile Thr Gly Arg Val Glu Asp
                          150
      (2) INFORMATION FOR SEQ ID NO:67:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 117 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..117
                (D) OTHER INFORMATION: / Ceres Seq. ID 1026564
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:
     Met Ala Leu Ile Ile Thr Cys Ser Ala Leu Pro Thr Ile Arg Ala Ser
                                         10
      Ser Gly Ser Gly Ser Leu Asn Pro Asp Gln Asn Arg Lys Lys Ser Ala
                                      25
55
      Ala Trp Trp Ala Pro Leu Phe Gly Leu Pro Ser Asp Pro Asp Tyr Leu
                                  40
      Asn Ile Glu Ser Ser Cys Ser Thr Val Asn Pro Asp Lys Thr Asp Ile
                              55
      Ser Gly Ser Gly Gln Lys Phe Arg Arg Gly Cys Phe Thr Glu Glu Lys
60
                          70
                                              75
```

Ala Lys Gln Leu Arg Arg Lys Thr Ala Glu Ala Ser Thr Phe His Asp

```
Val Met Tyr His Ser Ala Ile Ala Ser Arg Leu Ala Ser Asp Ile Thr
                100 105
      Gly Arg Val Glu Asp
             115
 5
      (2) INFORMATION FOR SEQ ID NO:68:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 484 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
10
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
               (B) LOCATION: 1..484
15
               (D) OTHER INFORMATION: / Ceres Seq. ID 1026648
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
     aaggatcaca aaattagggt tttaatttga tagagaagat gaatttcaga agctttgagg
      agttctggcc tttctacatg atgcaacact cgaatccatc gacgcgaagg ttgcacttca
                                                                           120
      taggtateat egegageate gttgetttga tatgttegat tttgateaac tggtggttet
                                                                           180
20
      tagetetggt gestetgett gggtaeggat tegegtggta tagesactte ttegtggaag
                                                                           240
      ggaatgttcc ggcgagcttt gggcatccgc tttggtcgtt tctctgcgat ctcaaqatgt
                                                                           300
      ttagtctgat gctcacagga agcatggaga gagagatgaa gagacttggt aagaggccat
                                                                           360
      tgttgcagct ctcttgaagc agcagcagat atctctatag aattgttctt gattcttctt
                                                                           420
      attgcqttct qcqattqqat tttaqactcc aqtttqtaat tacttcatqq aatcqttqtt
                                                                           480
25
     tgcg
      (2) INFORMATION FOR SEO ID NO:69:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 112 amino acids
                (B) TYPE: amino acid
30
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..112
35
                (D) OTHER INFORMATION: / Ceres Seq. ID 1026649
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:
     Met Asn Phe Arg Ser Phe Glu Glu Phe Trp Pro Phe Tyr Met Met Gln
               5
                                       10
40
      His Ser Asn Pro Ser Thr Arg Arg Leu His Phe Ile Gly Ile Ile Ala
                                    25
                  20
      Ser Ile Val Ala Leu Ile Cys Ser Ile Leu Ile Asn Trp Trp Phe Leu
                                 40
      Ala Leu Val Pro Leu Leu Gly Tyr Gly Phe Ala Trp Tyr Ser His Phe
45
                             55
      Phe Val Glu Gly Asn Val Pro Ala Ser Phe Gly His Pro Leu Trp Ser
                         70
                                             75
      Phe Leu Cys Asp Leu Lys Met Phe Ser Leu Met Leu Thr Gly Ser Met
                    85
                                         90
50
      Glu Arg Glu Met Lys Arg Leu Gly Lys Arg Pro Leu Leu Gln Leu Ser
                                     105
      (2) INFORMATION FOR SEQ ID NO:70:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 99 amino acids
55
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
60
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..99
                (D) OTHER INFORMATION: / Ceres Seq. ID 1026650
```

626 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:70: Met Met Gln His Ser Asn Pro Ser Thr Arg Arg Leu His Phe Ile Gly 10 Ile Ile Ala Ser Ile Val Ala Leu Ile Cys Ser Ile Leu Ile Asn Trp 5 25 Trp Phe Leu Ala Leu Val Pro Leu Leu Gly Tyr Gly Phe Ala Trp Tyr 40 Ser His Phe Phe Val Glu Gly Asn Val Pro Ala Ser Phe Gly His Pro 55 10 Leu Trp Ser Phe Leu Cys Asp Leu Lys Met Phe Ser Leu Met Leu Thr 70 75 Gly Ser Met Glu Arg Glu Met Lys Arg Leu Gly Lys Arg Pro Leu Leu 85 90 Gln Leu Ser 15 (2) INFORMATION FOR SEQ ID NO:71: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 98 amino acids (B) TYPE: amino acid 20 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 25 (B) LOCATION: 1..98 (D) OTHER INFORMATION: / Ceres Seq. ID 1026651 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71: Met Gln His Ser Asn Pro Ser Thr Arg Arg Leu His Phe Ile Gly Ile 5 10 30 Ile Ala Ser Ile Val Ala Leu Ile Cys Ser Ile Leu Ile Asn Trp Trp 25 Phe Leu Ala Leu Val Pro Leu Leu Gly Tyr Gly Phe Ala Trp Tyr Ser 40 His Phe Phe Val Glu Gly Asn Val Pro Ala Ser Phe Gly His Pro Leu 35 55 Trp Ser Phe Leu Cys Asp Leu Lys Met Phe Ser Leu Met Leu Thr Gly 70 75 Ser Met Glu Arg Glu Met Lys Arg Leu Gly Lys Arg Pro Leu Leu Gln 90 40 Leu Ser (2) INFORMATION FOR SEQ ID NO:72: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 563 base pairs 45 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: 50 (A) NAME/KEY: -(B) LOCATION: 1..563 (D) OTHER INFORMATION: / Ceres Seq. ID 1027881 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:72: aaaacacaga gagtgacgag agagagagag tgaaaatgga gaagtacttt ggaaatgcgt 60 55 120 acaggggaga tocaggagtg ccacatgcag atgcagatcg tttcatgaat atatggattg gttctgctgc tttctccgtt cttacctggg ttaatcctta catgtggcag ctctctaacc 180 agttcaatta ccatgacaag tggatgctgt ttgagcagta ccactggaaa aaagcaaggg 240 caaagaagca accttatgaa ttcaagtgga ataagatacc caaagaagtc agggactcgt 300 actattacaa ctggccagtc tacttcccat agaagtgtct ctgtgttgct gtgaatcaga 360

agagaaacca aacacttgtg gaacttctat ggaatctatc tctatctctt cttgatctgt

tattcgtage acttttggat tttaagtttt ttcttttgta atgatcaatc tctcaactct

teactgttte ateteeteta gttgaetete ttatggaaat attgaatete tttgaagage

420

480 5**4**0

```
agaaccagaa tctgtctgtt atg
      (2) INFORMATION FOR SEQ ID NO:73:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 109 amino acids
 5
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
10
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..109
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027882
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:
      Asn Thr Glu Ser Asp Glu Arg Glu Arg Val Lys Met Glu Lys Tyr Phe
15
                                          10
      Gly Asn Ala Tyr Arg Gly Asp Pro Gly Val Pro His Ala Asp Ala Asp
                                      25
                  20
                                                          3.0
      Arg Phe Met Asn Ile Trp Ile Gly Ser Ala Ala Phe Ser Val Leu Thr
                                  4.0
20
      Trp Val Asn Pro Tyr Met Trp Gln Leu Ser Asn Gln Phe Asn Tyr His
                              55
                                                  60
      Asp Lys Trp Met Leu Phe Glu Gln Tyr His Trp Lys Lys Ala Arg Ala
                          70
                                              75
      Lys Lys Gln Pro Tyr Glu Phe Lys Trp Asn Lys Ile Pro Lys Glu Val
25
                     85
                                          90
      Arg Asp Ser Tyr Tyr Asn Trp Pro Val Tyr Phe Pro
                  100
      (2) INFORMATION FOR SEQ ID NO:74:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 98 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..98
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027883
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:74:
40
      Met Glu Lys Tyr Phe Gly Asn Ala Tyr Arg Gly Asp Pro Gly Val Pro
                  5
                                          10
                                                              15
      His Ala Asp Ala Asp Arg Phe Met Asn Ile Trp Ile Gly Ser Ala Ala
                                      25
      Phe Ser Val Leu Thr Trp Val Asn Pro Tyr Met Trp Gln Leu Ser Asn
45
                                  40
      Gln Phe Asn Tyr His Asp Lys Trp Met Leu Phe Glu Gln Tyr His Trp
                              55
      Lys Lys Ala Arg Ala Lys Lys Gln Pro Tyr Glu Phe Lys Trp Asn Lys
                          70
                                              75
50
      Ile Pro Lys Glu Val Arg Asp Ser Tyr Tyr Tyr Asn Trp Pro Val Tyr
                                          90
                      85
      Phe Pro
      (2) INFORMATION FOR SEQ ID NO:75:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 75 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
```

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628
                (B) LOCATION: 1..75
                (D) OTHER INFORMATION: / Ceres Seq. ID 1027884
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:75:
      Met Asn Ile Trp Ile Gly Ser Ala Ala Phe Ser Val Leu Thr Trp Val
 5
                                          10
      Asn Pro Tyr Met Trp Gln Leu Ser Asn Gln Phe Asn Tyr His Asp Lys
                  20
                                      25
      Trp Met Leu Phe Glu Gln Tyr His Trp Lys Lys Ala Arg Ala Lys Lys
                                  40
                                                       4.5
10
      Gln Pro Tyr Glu Phe Lys Trp Asn Lys Ile Pro Lys Glu Val Arg Asp
                              55
                                                  60
      Ser Tyr Tyr Asn Trp Pro Val Tyr Phe Pro
                          70
      (2) INFORMATION FOR SEQ ID NO:76:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 305 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
20
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..305
                (D) OTHER INFORMATION: / Ceres Seq. ID 1381797
25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:76:
      gaggttttgg ttttgttgac tcagtgagga ttttctaggc gattttctcg agaaaatcgt
      ttgaggaaaa tggagacttc tatgaggtat accagcaatt ccaagtctat gaagattcat
      gccaaagaga aggttccggt gaactcaaaa acccatttac agcttcatgg agagttagat
      actggaactg gggctccgag ttacttctgt gcgatgatta gacacttttt tcctgaggct
30
      tcaacaggcc ttggggtagg attgcattat gataagcgcc aaaagcttcg gtgtcttgta
      cgcgg
      (2) INFORMATION FOR SEQ ID NO:77:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 78 amino acids
35
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..78
                (D) OTHER INFORMATION: / Ceres Seq. ID 1381798
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77:
      Met Glu Thr Ser Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile
45
                      5
                                           1.0
      His Ala Lys Glu Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu
      His Gly Glu Leu Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala
                                  40
                                                       45
50
      Met Ile Arg His Phe Phe Pro Glu Ala Ser Thr Gly Leu Gly Val Gly
                              55
      Leu His Tyr Asp Lys Arg Gln Lys Leu Arg Cys Leu Val Arg
                          70
      (2) INFORMATION FOR SEQ ID NO:78:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
```

(A) NAME/KEY: peptide

629

```
(B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 1381799
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:78:
     Met Arg Tyr Thr Ser Asn Ser Lys Ser Met Lys Ile His Ala Lys Glu
 5
                                          10
     Lys Val Pro Val Asn Ser Lys Thr His Leu Gln Leu His Gly Glu Leu
                  20
                                      25
     Asp Thr Gly Thr Gly Ala Pro Ser Tyr Phe Cys Ala Met Ile Arg His
                                  40
                                                      45
10
      Phe Phe Pro Glu Ala Ser Thr Gly Leu Gly Val Gly Leu His Tyr Asp
                              55
                                                  60
     Lys Arg Gln Lys Leu Arg Cys Leu Val Arg
                         70
      (2) INFORMATION FOR SEQ ID NO:79:
15
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 65 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
20
          /ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 1381800
25
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:
     Met Lys Ile His Ala Lys Glu Lys Val Pro Val Asn Ser Lys Thr His
                                         10
      Leu Gln Leu His Gly Glu Leu Asp Thr Gly Thr Gly Ala Pro Ser Tyr
                  20
                                      25
30
      Phe Cys Ala Met Ile Arg His Phe Phe Pro Glu Ala Ser Thr Gly Leu
                                  40
      Gly Val Gly Leu His Tyr Asp Lys Arg G!n Lys Leu Arg Cys Leu Val
                              55
      Arg
35
      65
      (2) INFORMATION FOR SEQ ID NO:80:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 840 base pairs
                (B) TYPE: nucleic acid
40
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
45
                (B) LOCATION: 1..840
                (D) OTHER INFORMATION: / Ceres Seq. ID 1442747
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:
      acacgaaacg attccgcttc tccgcttcct tcataaatat ctcactcatc gtcactacat
      tgttctttca cacgcaattt tcatttctct cttaaccgta aacgaatctc tcttctcaaa
                                                                             120
50
      gttttatttt ctctctgcga tggctcaaga agatgttact gctgttgcta ctaacggtgc
                                                                             180
      tggtccggtg gagacgcatc ttgtcttcac tgagttcaag cagatgttgc tcgttgaagc
                                                                             240
      tcaaaaggtc ggtgacgctg ttactttcta caaatctgct tttggtgcga tcgagtctgg
                                                                             300
      tcattctctt taccctaagc gtaacttgac caagagcttc ctcatgttct ctcttctgag
                                                                             360
      cttaatctcg ctggctcttc cttcgttgtt tgcgacgttt cctctcccc tggtttttct
                                                                             420
55
      actgcgaaat cggaaggttc gggagtgact tttcttctcg gaactaagga tgctgaagcc
                                                                             480
                                                                             540
      gccgttgcga aagctgttga cgccggagct gtgaaagtgg aggttacgga ggcagaagtt
      gaactgggat tcaaaggaaa agttacggat ccttttggtg tcacttggat cttcgcggag
                                                                             600
      aagaagaccg tgatcaccga cgagaacaaa gaggtttaga atctgtcgtc ggatctaatt
                                                                             660
      cttcgtcggt ttctgaacaa aaaaaaatta tctattatcc taggtttatg cttttatttt
                                                                             720
60
                                                                            780
      gtcttttgtg gaatccggta atcagtaaac cggatcgagt aggctaatga ctttcggatt
      ctaatttcac cccttttgac aaactctaat cttggtgaaa ttgcatatta atctcgggct
                                                                             840
```

(2) INFORMATION FOR SEQ ID NO:81:

```
(i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 148 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
 5
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (P) LOCATION: 1..148
10
                (D) OTHER INFORMATION: / Ceres Seq. ID 1442748
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:81:
     His Glu Thr Ile Pro Leu Leu Arg Phe Leu His Lys Tyr Leu Thr His
                     5
                                         10
     Arg His Tyr Ile Val Leu Ser His Ala Ile Phe Ile Ser Leu Leu Thr
15
                 20
                                      25
      Val Asn Glu Ser Leu Phe Ser Lys Phe Tyr Phe Leu Ser Ala Met Ala
                                  40
      Gln Glu Asp Val Thr Ala Val Ala Thr Asn Gly Ala Gly Pro Val Glu
                              5.5
2.0
      Thr His Leu Val Phe Thr Glu Phe Lys Gln Met Leu Leu Val Glu Ala
                          70
                                              75
      Gln Lys Val Gly Asp Ala Val Thr Phe Tyr Lys Ser Ala Phe Gly Ala
                      85
                                          90
      Ile Glu Ser Gly His Ser Leu Tyr Pro Lys Arg Asn Leu Thr Lys Ser
25
                  100
                                      105
                                                         110
      Phe Leu Met Phe Ser Leu Leu Ser Leu Ile Ser Leu Ala Leu Pro Ser
                                  120
      Leu Fhe Ala Thr Phe Pro Leu Ser Leu Val Phe Leu Leu Arg Asn Arg
         130
                              135
                                                  140
30
      Lys Val Arg Glu
      145
      (2) INFORMATION FOR SEQ IT NO:82:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 102 amino acids
35
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          ix) FEATURE:
40
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..102
                (D) OTHER INFORMATION: / Ceres Seq. ID 1442749
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:82:
      Met Ala Gln Glu Asp Val Thr Ala Val Ala Thr Asn Gly Ala Gly Pro
45
                  5
                                                             15
                                         10
      Val Glu Thr His Leu Val Phe Thr Glu Phe Lys Gln Met Leu Leu Val
                                      25
      Glu Ala Gln Lys Val Gly Asp Ala Val Thr Phe Tyr Lys Ser Ala Phe
                                                     45
                                 40
      Gly Ala Ile Glu Ser Gly His Ser Leu Tyr Pro Lys Arg Asn Leu Thr
50
                              55
                                              60
      Lys Ser Phe Leu Met Phe Ser Leu Leu Ser Leu Ile Ser Leu Ala Leu
                          70
                                          75
      Pro Ser Leu Phe Ala Thr Phe Pro Leu Ser Leu Val Phe Leu Leu Arg
55
                     85
                                          90
      Asn Arg Lys Val Arg Glu
                 100
      (2) INFORMATION FOR SEQ ID NO:83:
           (i) SEQUENCE CHARACTERISTICS:
60
                (A) LENGTH: 74 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
```

631

Pro Ser Ser

```
(D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
 5
                (B) LOCATION: 1..74
                (D) OTHER INFORMATION: / Ceres Seq. ID 1442750
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:83:
      Met Leu Leu Val Glu Ala Gln Lys Val Gly Asp Ala Val Thr Phe Tyr
                                          10
10
      Lys Ser Ala Phe Gly Ala Ile Glu Ser Gly His Ser Leu Tyr Pro Lys
                  20
                                                           30
                                      25
      Arg Asn Leu Thr Lys Ser Phe Leu Met Phe Ser Leu Leu Ser Leu Ile
                                  40
                                                      45
      Ser Leu Ala Leu Pro Ser Leu Phe Ala Thr Phe Pro Leu Ser Leu Val
15
                              55
                                                   60
      Phe Leu Leu Arg Asn Arg Lys Val Arg Glu
                          70
      (2) INFORMATION FOR SEQ ID NO:84:
           (i) SEQUENCE CHARACTERISTICS:
20
                A) LENGTH: 513 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
25
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..513
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459199
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:84:
30
      aacttggatc ttaacacaga gaagaagcac aatcggaaga aaggagaaga cgaagatgtc
      gttggtatgg ctggaagcga tgttgcctct cggaatcatc ggtgggatgc tctgtatcat
                                                                             120
      gggcaattot cagtactaca tocacaaago ttatcatggo cgtoctaago acatoggoca
                                                                             180
      cgatgaatgg gatgttgcta tggaaagacg cgacaagaaa gtcgtcgaga aagctgcagc
                                                                             240
      teetteetea tgattegett tatetetttt gtgtteetea ggtgaceaet tgtggtgaca
                                                                             300
35
      aataaagtgc attccagaag aagaagaagc tgggggatct agtactttca ttcccatttg
                                                                             360
      attttccttg qacatattaa agctttcaqa aatcaqacct caataacatt tggtttatca
                                                                             420
      atatttctct attcgtgata ttttgtatgc ttttaagttg catgatacac tgaactactc
                                                                             480
      tottgctgtg tgagtgaata aatgaatato tgo
      (2) INFORMATION FOR SEQ ID NO:85:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 83 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..83
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459200
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:85:
      Thr Trp Ile Leu Thr Gln Arg Arg Ser Thr Ile Gly Arg Lys Glu Lys
                                           10
      Thr Lys Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile
                                       25
                  2.0
55
      Ile Gly Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His
                                   40
      Lys Ala Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp
                              55
                                                   60
      Val Ala Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Ala
60
                          70
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6

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632
      (2) INFORMATION FOR SEQ ID NO:86:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 65 amino acids
                (B) TYPE: amino acid
 5
                (C) STRANDEDNESS:
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
10
               (B) LOCATION: 1..65
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459201
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:86:
     Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile Ile Gly
                                         10
15
     Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His Lys Ala
                 20
                                      25
      Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp Val Ala
                                 4.0
     Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Pro Ser
20
                             55
                                                60
      Ser
      65
      (2) INFORMATION FOR SEQ ID NO:87:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 57 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..57
                (D) OTHER INFORMATION: / Ceres Seq. ID 1459202
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:87:
35
      Met Leu Pro Leu Gly Ile Ile Gly Gly Met Leu Cys Ile Met Gly Asn
                                         10
      Ser Gln Tyr Tyr Ile His Lys Ala Tyr His Gly Arg Pro Lys His Ile
                              25 30
                 20
      Gly His Asp Glu Trp Asp Val Ala Met Glu Arg Arg Asp Lys Lys Val
40
             35
                                40
      Val Glu Lys Ala Ala Ala Pro Ser Ser
                             55
      (2) INFORMATION FOR SEQ ID NO:88:
           (i) SEQUENCE CHARACTERISTICS:
45
                (A) LENGTH: 1140 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
50
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1140
                (D) OTHER INFORMATION: / Ceres Seq. ID 1565605
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:88:
55
      atqatttqtt tacatttttt cqtqqtqttc ttqatttttt tctttqatca aaactaqtaq
                                                                            60
      aaattttcca atttcgtcct gccttgtgat tagtagtaga tattttggaa tttgattagt
                                                                            120
      aggtagaaga agatggggac gacgttagat gtatcaagag cagagctagc acttgtggta
                                                                           180
      atgtatttga acaaagcaga ggcaagagat aagttatgca gagctataca gtatggttcc
      aagttettga gtggtggaca acetggtaet geteaaaatg ttgacaaate tactagetta
60
      gcaagaaaag tottoogtot tittoaagtit gtgaatgact tgcatggtot tatoagtoot
                                                                            360
      gtgcctaaag gaactcctct tcctcttgtt ttacttggaa agtcgaagaa cgcactttta
                                                                           420
```

tctacattct tgttcctgga tcaaattgtc tggcttggga gatcaggaat atataagaac

633 aaagaacgag ctgagttact tggacgtata tctctcttct gctggatggg atcttctgtc 540 tgcacaactt tagtcgaggt tggtgagatg ggaaggcttt cttcatcaat gaagaagatc 600 gaaaagggac tcaagaatgg aaacaagtat caggatgagg attatcgtgc taagctaaaa 660 aaatcaaacg agaggtcact tgctttgatc aaatcagcta tggacattgt tgtagcagct 720 5 ggtcttcttc agttagctcc aacgaagatc actcctcqtg tcaccggagc ttttggattc 780 atcacctcca tcatttcttg ttaccagttg cttccgacac gccccaagat caaaacaccc 840 900 tgaagtcaat cgaggaagct ggtgttaagg agaaagtcat ttcagaaagt ttctgttctt cttattatta aggaaattca ctgctgttat aaaacatcgt ataaaatact tatttgcagg 960 1020 aattacggtt attataaatt tatgcctcag attcttgata cacaagtttc aatactcaat 10 agtattgcat tattatgaga tgttgtttct Gctacgcaat aactagaggc aaaatccaac 1080 1140 tacatatatg ttttgatctg attattataa tagactttgg gcttcattcc gcctaaagct (2) INFORMATION FOR SEQ ID NO:89: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 236 amino acids 15 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 20 (A) NAME/KEY: peptide (B) LOCATION: 1..236 (D) OTHER INFORMATION: / Ceres Seq. ID 1565606 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:89: Met Gly Thr Thr Leu Asp Val Ser Arg Ala Glu Leu Ala Leu Val Val 25 1.0 Met Tyr Leu Asn Lys Ala Glu Ala Arg Asp Lys Leu Cys Arg Ala Ile 20 25 30 Gln Tyr Gly Ser Lys Phe Leu Ser Gly Gly Gln Pro Gly Thr Ala Gln 40 30 Asn Val Asp Lys Ser Thr Ser Leu Ala Arg Lys Val Phe Arg Leu Phe 55 Lys Phe Val Asn Asp Leu His Gly Leu Ile Ser Pro Val Pro Lys Gly 70 75 Thr Pro Leu Pro Leu Val Leu Leu Gly Lys Ser Lys Asn Ala Leu Leu 35 Ser Thr Phe Leu Phe Leu Asp Gln Ile Val Trp Leu Glv Arg Ser Gly 105 110 Ile Tyr Lys Asn Lys Glu Arg Ala Glu Leu Leu Glv Arg Ile Ser Leu 120 125 40 Phe Cys Trp Met Gly Ser Ser Val Cys Thr Thr Leu Val Glu Val Gly 135 140 Glu Met Gly Arg Leu Ser Ser Met Lys Lys Ile Glu Lys Gly Leu 150 155 Lys Asn Gly Asn Lys Tyr Gln Asp Glu Asp Tyr Arg Ala Lys Leu Lys 45 170 165 175 Lys Ser Asn Glu Arg Ser Leu Ala Leu Ile Lys Ser Ala Met Asp Ile 180 185 Val Val Ala Ala Gly Leu Leu Gln Leu Ala Pro Thr Lys Ile Thr Pro 200 205 50 Arg Val Thr Gly Ala Phe Gly Phe Ile Thr Ser Ile Ile Ser Cys Tyr 215 Gln Leu Leu Pro Thr Arg Pro Lys Ile Lys Thr Pro 230 (2) INFORMATION FOR SEQ ID NO:90: 55 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 220 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 60 (ii) MOLECULE TYPE: peptide (ix) FEATURE:

(A) NAME/KEY: peptide

```
(B) LOCATION: 1..220
                (D) OTHER INFORMATION: / Ceres Seq. ID 1565607
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:90:
     Met Tyr Leu Asn Lys Ala Glu Ala Arg Asp Lys Leu Cys Arg Ala Ile
5
                      5
                                          10
      Gln Tyr Gly Ser Lys Phe Leù Ser Glv Gly Gln Pro Gly Thr Ala Gln
                                      25
     Asn Val Asp Lys Ser Thr Ser Leu Ala Arg Lys Val Phe Arg Leu Phe
10
     Lys Phe Val Asn Asp Leu His Gly Leu Ile Ser Pro Val Pro Lys Gly
                              55
      Thr Pro Leu Pro Leu Val Leu Leu Gly Lys Ser Lys Asn Ala Leu Leu
                          70
                                               75
      Ser Thr Phe Leu Phe Leu Asp Gln Ile Val Trp Leu Gly Arg Ser Gly
15
                                           90
      Ile Tyr Lys Asn Lys Glu Arg Ala Glu Leu Leu Gly Arg Ile Ser Leu
                                      105
                  100
                                                           110
      Phe Cys Trp Met Gly Ser Ser Val Cys Thr Thr Leu Val Glu Val Gly
                                  120
                                                       125
              115
20
      Glu Met Gly Arg Leu Ser Ser Ser Met Lys Lys Ile Glu Lys Gly Leu
                              135
                                                   140
      Lys Asn Gly Asn Lys Tyr Gln Asp Glu Asp Tyr Arg Ala Lys Leu Lys
      145
                          150
                                               155
      Lvs Ser Asn Glu Arg Ser Leu Ala Leu Ile Lys Ser Ala Met Asp Ile
25
                      165
                                           170
      Val Val Ala Ala Gly Leu Leu Gln Leu Ala Pro Thr Lys Ile Thr Pro
                                      185
      Arg Val Thr Gly Ala Phe Gly Phe Ile Thr Ser Ile Ile Ser Cys Tyr
                                  200
30
      Gln Leu Leu Pro Thr Arg Pro Lys Ile Lys Thr Pro
                              215
      (2) INFORMATION FOR SEQ ID NO:91:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 533 base pairs
35
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
40
                (A) NAME/KEY: -
                (B) LOCATION: 1..533
                (D) OTHER INFORMATION: / Ceres Seq. ID 1566686
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:91:
                                                                              60
      atgattggtg gtttcttcgc cgatgttaaa ttcaacagat cgagagagaa aaaactctct
45
                                                                             120
      ctccctcctc cgttcaccgt cgcgactcgg ctctacagta cattttgttg agcaatctcg
                                                                             180
      tatttagagg attccaatgt ctggcgcaga ggataataaa tcttcccatg ctcagctttc
      gtcccaaatt tttcttgatc ttgtggattc ggtgatagct gatgtagcat ctgagtgtca
                                                                             240
                                                                             300
      tcgggtagca cgactaggac ttgatcgtga tttggatata gtagaagaag agttgagatt
                                                                             360
      gtcagtggaa gcgcgtgcga agatcgctga tcctagcaac aacctcgaaa ccaacaccaa
50
                                                                             420
      atatgttgtt gatatatttg ggcagactca ccctcctgta gctagtgaag tgttcaattg
      catgaactgt qgqcqacaaa ttgttqcaqq aaqgtttqct cctcatctgg aaaaatgcat
                                                                             480
      gggaaaggga agaaaggctc gtgGccaagG actaccagaa gcacaacggc tgc
      (2) INFORMATION FOR SEQ ID NO:92:
           (i) SEQUENCE CHARACTERISTICS:
55
                (A) LENGTH: 132 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
60
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..132
```

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(D) OTHER INFORMATION: / Ceres Seq. ID 1566687
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:92:
     Met Ser Gly Ala Glu Asp Asn Lys Ser Ser His Ala Gln Leu Ser Ser
                                         10
 5
     Gln Ile Phe Leu Asp Leu Val Asp Ser Val Ile Ala Asp Val Ala Ser
                                     25
     Glu Cys His Arg Val Ala Arg Leu Gly Leu Asp Arg Asp Leu Asp Ile
                                 4.0
     Val Glu Glu Leu Arg Leu Ser Val Glu Ala Arg Ala Lys Ile Ala
10
                             55
     Asp Pro Ser Asn Asn Leu Glu Thr Asn Thr Lys Tyr Val Val Asp Ile
                         70
                                             75
     Phe Gly Gln Thr His Pro Pro Val Ala Ser Glu Val Phe Asn Cys Met
                                         90
15
     Asn Cys Gly Arg Gln Ile Val Ala Gly Arg Phe Ala Pro His Leu Glu
                 100
                                     105
     Lys Cys Met Gly Lys Gly Arg Lys Ala Arg Gly Gln Gly Leu Pro Glu
                                 120
                                                     125
     Ala Gln Arg Leu
20
          130
      (2) INFORMATION FOR SEQ ID NO:93:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 815 base pairs
                (B) TYPE: nucleic acid
25
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
30
                (B) LOCATION: 1..815
                (D) OTHER INFORMATION: / Ceres Seq. ID 1567367
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:93:
      60
                                                                           120
      ttcaatggct gtcgtcggcg ctccaatatc gtctccggcg gctcagctgc agacacaatt
35
                                                                           180
      tototocaat occattotoo occgotttog ocgqtottto tocacoggaa aatoaccago
                                                                           240
      aactttctcc gtcgtagcta tggctcccca gaaaaaggtg aacaaatatg atgccaagtg
      gaaqaaacaa tqqtacqqaq ctqqattqtt tttcqaaqqq aqtqaqcaaa taaacqttqa
                                                                           300
      tgttttcaag aagctggaga agcgaaaagt gttgagcaac gttgagaaat ctggcctgct
                                                                           360
      qtaaaaqcaq aqqqqttqqq actcacattq tcatctcttq aqaaqcttaa aqtcttctcc
                                                                          420
40
                                                                          480
      aaaqcaqaqq accttqqtct tctcaqtctc cttqaqaact taqctqqamG AAcatcqcct
                                                                          540
      geggtettag ceteggetge attaceaget etcaeggetg etattgtage egtggtgttg
      atcccqaatq actcaactac tctagtggtt gctcaggcgg ttttggccqg tgctcttgcg
                                                                           600
      cttacagggg ttgttttgtt ggttggttct gttgttttgg atggacttca agaagctgac
                                                                           660
      tgattctttc tctgtaaacc aaacataaac ccatgtcttg tccaattgat ttttgtcagt
                                                                          720
45
      tgctgattta tagctgtatg gttcagttgt ttatggttag tccaagacat aagctgagtg
                                                                          780
      atagaaagaa gctttataat aattaaacaa atatt
      (2) INFORMATION FOR SEQ ID NO:94:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 76 amino acids
50
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
55
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..76
                (D) OTHER INFORMATION: / Ceres Seq. ID 1567368
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:94:
      Glu Thr Arg His Glu Arg Ile Ser Lys Glu Ile Ile Phe Ile Ala Ser
60
                                         10
      Leu Arg Phe Phe Asn Gly Cys Arg Arg Arg Ser Asn Ile Val Ser
                  20
                                      25
```

PCT/US00/00466

WO 00/40695 636 Gly Gly Ser Ala Ala Asp Thr Ile Ser Leu Gln Ser His Ser Pro Pro 40 Leu Ser Pro Val Phe Leu His Arg Lys Ile Thr Ser Asn Phe Leu Arg 55 Arg Ser Tyr Gly Ser Pro Glu Lys Gly Glu Gln Ile 70 (2) INFORMATION FOR SEQ ID NO:95: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 120 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPCLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..120 (D) OTHER INFORMATION: / Ceres Seq. ID 1567369 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:95: Lys Gln Asp Thr Lys Glu Ser Leu Lys Lys Ser Phe Ser Ser Leu Arg 5 10 Phe Asp Phe Ser Ser Met Ala Val Val Gly Ala Pro Ile Ser Ser Pro 20 25 Ala Ala Gln Leu Gln Thr Gln Phe Leu Ser Asn Pro Ile Leu Pro Arg 40 Phe Arg Arg Ser Phe Ser Thr Gly Lys Ser Pro Ala Thr Phe Ser Val 55 Val Ala Met Ala Pro Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys Trp 70 75 Lys Lys Gln Trp Tyr Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu Gln 85 90 Ile Asn Val Asp Val Phe Lys Lys Leu Glu Lys Arg Lys Val Leu Ser 100 105 110

30

Asn Val Glu Lys Ser Gly Leu Leu 115

- 35 (2) INFORMATION FOR SEQ ID NO:96:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 99 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS:
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:

5

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- (A) NAME/KEY: peptide
- (B) LOCATION: 1..99
- 45 (D) OTHER INFORMATION: / Ceres Seq. ID 1567370

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:96:

Met Ala Val Val Gly Ala Pro Ile Ser Ser Pro Ala Ala Gln Leu Gln 10

Thr Gln Phe Leu Ser Asn Pro Ile Leu Pro Arg Phe Arg Arg Ser Phe 50 20 25

Ser Thr Gly Lys Ser Pro Ala Thr Phe Ser Val Val Ala Met Ala Pro 40

Gln Lys Lys Val Asn Lys Tyr Asp Ala Lys Trp Lys Lys Gln Trp Tyr 55

55 Gly Ala Gly Leu Phe Phe Glu Gly Ser Glu Gln Ile Asn Val Asp Val 70 75

Phe Lys Lys Leu Glu Lys Arg Lys Val Leu Ser Asn Val Glu Lys Ser 90

Gly Leu Leu

(2) INFORMATION FOR SEQ ID NO:97:

(i) SEQUENCE CHARACTERISTICS:

```
637
                (A) LENGTH: 541 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
 5
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..541
                (D) OTHER INFORMATION: / Ceres Seq. ID 1570101
10
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:97:
     aaaMctttac qttqqtatta tctqqtttca cattttctta accctctttq qtqaatqcqt
                                                                             60
      cattgaaacg ccattatccg ccgtgctaat ctgatctcct cccccgtcgg aaaaaatggc
                                                                             120
      cactggtgga aaagtotoot toaaagtoac totgacttog gatootaago ttootttoaa
                                                                             180
      agttttcagc gtaccagaag gagctccgtt cacggcggtt ctgaaattcg cagcagaaga
                                                                             240
15
      gttcaaggtt cctccacaaa ccagegccat catcactaat gatgggatcg ggatcaatce
                                                                             300
      toaacagagt gcaggaaacg tttttctgaa gcacggatct gaactaagat tgatcccgcg
                                                                             360
      tgatagagtt ggagctgtgt ttgtgatgga tccatagatg atgtgtaagc aatgattaaa
                                                                             420
      aactcagttc caaaaaaaa acataataat atactgaaat gcttatgtat caactTcqtt
                                                                             480
      qaataaatat gtaatogtoa catootttta oqqattgaag tgagttgaat ttggtcaagt
                                                                            540
20
      (2) INFORMATION FOR SEQ ID NO: 98:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 93 amino acids
                (B) TYPE: amino acid
25
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
30
                (B) LOCATION: 1..93
                (D) OTHER INFORMATION: / Ceres Seq. ID 1570102
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:98:
      Met Ala Thr Gly Gly Lys Val Ser Phe Lys Val Thr Leu Thr Ser Asp
                                          10
      Pro Lys Leu Pro Phe Lys Val Phe Ser Val Pro Glu Gly Ala Pro Phe
3.5
                  20
                                      25
                                                           30
      Thr Ala Val Leu Lys Phe Ala Ala Glu Glu Phe Lys Val Pro Pro Gln
                                  40
                                                       4.5
      Thr Ser Ala Ile Ile Thr Asn Asp Gly Ile Gly Ile Asn Pro Gln Gln
40
                              55
                                                   60
      Ser Ala Gly Asn Val Phe Leu Lys His Gly Ser Glu Leu Arg Leu Ile
                          70
                                               75
      Pro Arg Asp Arg Val Gly Ala Val Phe Val Met Asp Pro
                      85
45
      (2) INFORMATION FOR SEQ ID NO:99:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 418 base pairs
                 (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
50
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
                 (B) LOCATION: 1..418
5.5
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571051
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:99:
      aaaaggaaga cgacgatgca gagtttaacg tcaaacgaag tTgcgggttt ggcagttggg
                                                                              60
      gctctgcttc ttggtgccac categctgct cctaaagttg atgcttttat cgctgcttct
                                                                             120
      cagagaagat ctcttggcat gtgtcgaaaa tgtggagatc ttaagaatgt agcttgtggc
                                                                             180
```

cgttgcaaag gaacaggaac aatcaaatca ggaggattct ttggtttcag tgactcatca

aacacaagat cagtggcttg cgataattgc caagccaaag gttgtttccc ttgccctgaa

tgctcaaaat cttgaccatt ttctcggtat tttatagttg tttcatcttc ttgacactat

240

300

360

```
gataagtgta atcggtccat tggtaatggt aatgttaaag ttgaagaatg tcttgttt
      (2) INFORMATION FOR SEQ ID NO:100:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 104 amino acids
5
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          :ix) FEATURE:
10
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..104
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571052
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:100:
      Lys Arg Lys Thr Thr Met Gln Ser Leu Thr Ser Asn Glu Val Ala Gly
15
      1
                      5
                                          10
      Leu Ala Val Gly Ala Leu Leu Gly Ala Thr Ile Ala Ala Pro Lys
                  20
                                      25
      Val Asp Ala Phe Ile Ala Ala Ser Gln Arg Arg Ser Leu Gly Met Cys
                                                       4.5
20
      Arg Lys Cys Gly Asp Leu Lys Asn Val Ala Cys Gly Arg Cys Lys Gly
                              5.5
      Thr Gly Thr Ile Lys Ser Gly Gly Phe Phe Gly Phe Ser Asp Ser Ser
                                               75
                          70
      Asn Thr Arg Ser Val Ala Cys Asp Asn Cys Gln Ala Lys Gly Cys Phe
25
      Pro Cys Pro Glu Cys Ser Lys Ser
                 100
      (2) INFORMATION FOR SEQ ID NO:101:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 99 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..99
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571053
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:101:
40
      Met Gln Ser Leu Thr Ser Asn Glu Val Ala Gly Leu Ala Val Gly Ala
                                           10
      Leu Leu Gly Ala Thr Ile Ala Ala Pro Lys Val Asp Ala Phe Ile
                                       25
      Ala Ala Ser Gln Arg Arg Ser Leu Gly Met Cys Arg Lys Cys Gly Asp
45
                                   40
      Leu Lys Asn Val Ala Cys Gly Arg Cys Lys Gly Thr Gly Thr Ile Lys
                               55
      Ser Gly Gly Phe Phe Gly Phe Ser Asp Ser Ser Asn Thr Arg Ser Val
                          70
                                               75
50
      Ala Cys Asp Asn Cys Gln Ala Lys Gly Cys Phe Pro Cys Pro Glu Cys
                       85
                                           90
      Ser Lys Ser
      (2) INFORMATION FOR SEQ ID NO:102:
55
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 58 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
60
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
```

639

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(B) LOCATION: 1..58
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571054
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:102:
     Met Cys Arg Lys Cys Gly Asp Leu Lys Asn Val Ala Cys Gly Arg Cys
 5
                                          10
      Lys Gly Thr Gly Thr Ile Lys Ser Gly Gly Phe Phe Gly Phe Ser Asp
                                      25
                  20
      Ser Ser Asn Thr Arg Ser Val Ala Cys Asp Asn Cys Gln Ala Lys Gly
                                  40
10
      Cys Phe Pro Cys Pro Glu Cys Ser Lys Ser
          50
      (2) INFORMATION FOR SEQ ID NO:103:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 444 base pairs
15
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
20
                (A) NAME/KEY: -
                (B) LOCATION: 1..444
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571100
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:103:
      aatctacttt tgagttetet ggageagett tggtetttte tetetteagt tacagetege
                                                                              60
25
      gatccagatc catatccagt catggcgggt gtagtagcaa gaattgtgga aaagtttggt
                                                                             120
      aaacgtgata tttggtatcg aatatcgttg atatcaattc cagtctctag tgtcctaata
                                                                             180
      gtactaggga atatggagct tgctcgtatc agagaggaga agaagctttt gaaggagagg
                                                                             240
                                                                             300
      tttgatcaac tgagggctag aggaataatc agagatcagt aatccaaaag tgtgtttagg
                                                                             360
      ctttgttttt ttgtttcttt tctatgaatt taatttttta tttctccttt ccatgaatta
30
      accaactcta tgtaacgaga tgatttattc ttgtctctct aagactcgag atgattcgct
                                                                             420
      attgataaaa cttctattat tScc
      (2) INFORMATION FOR SEQ ID NO:104.
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 93 amino acids
35
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
40
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..93
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571101
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:104:
      Asn Leu Leu Ser Ser Leu Glu Gln Leu Trp Ser Phe Leu Ser Ser
45
                                           10
      Val Thr Ala Arg Asp Pro Asp Pro Tyr Pro Val Met Ala Gly Val Val
                                       25
      Ala Arg Ile Val Glu Lys Phe Gly Lys Arg Asp Ile Trp Tyr Arg Ile
                                   40
50
      Ser Leu Ile Ser Ile Pro Val Ser Ser Val Leu Ile Val Leu Gly Asn
                               55
                                                   60
      Met Glu Leu Ala Arg Ile Arg Glu Glu Lys Lys Leu Leu Lys Glu Arg
                           70
      Phe Asp Gln Leu Arg Ala Arg Gly Ile Ile Arg Asp Gln
55
                      85
       (2) INFORMATION FOR SEQ ID NO:105:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 66 amino acids
                 (B) TYPE: amino acid
60
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
```

(ii) MOLECULE TYPE: peptide

640

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(ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..66
                (D) OTHER INFORMATION: / Ceres Seq. ID 1571102
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:105:
     Met Ala Gly Val Val Ala Arq Ile Val Glu Lys Phe Gly Lys Arq Asp
                                          10
      Ile Trp Tyr Arg Ile Ser Leu Ile Ser Ile Pro Val Ser Ser Val Leu
                  20
                                      25
10
      Ile Val Leu Gly Asn Met Glu Leu Ala Arg Ile Arg Glu Glu Lys Lys
                                 40
      Leu Leu Lys Glu Arg Phe Asp Gln Leu Arg Ala Arg Gly Ile Ile Arg
          50
                             55
      Asp Gln
15
      65
      (2) INFORMATION FOR SEQ ID NO:106:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 636 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..636
                (D) OTHER INFORMATION: / Ceres Seq. ID 1665272
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:106:
      cagaataaaa atctatttct tgtgtttgtg ttatatttac ttaaaaaaaat aaaggagatc
                                                                             60
      agtttgggag aaagcagcaa agaagaaaaa tggcagcaac atttgcaaca ccatcgacgg
                                                                             120
30
      tgataggcct cggaggatca tccatcacca ccaaaccctt ctcttcatcc tttttaaaac
                                                                             180
      caacattaag cgccaagaac cctttgagac tcgccggtgc atcgggagga agagtcactt
                                                                             240
      gctttgagag gaactggttg aggagagatt tgaacgtggt aggatttggg ctgatcggat
                                                                             300
                                                                             360
      ggctagctcc gtcgagcatt ccagcgataa atgggaagag cctgacgggt ctcttcttcg
                                                                             420
      atagcatcgg aactgagete geteacttee egacteetee ageteteact teacagttet
35
      ggttgtggtt ggttacgtgg cacttaggcc tettectetg ceteacttte ggacaaateg
                                                                             480
      gattcaaggg caggactgag gattacttct aaggataact attcttgttt tcgtttgtac
                                                                             540
      tatatgctct ctcttggtta tgtgtaatat tatcaatcaa aacaaagctt tctttgctat
                                                                             600
      ttgatgtttc attctatatc tcgacaAgtc tttctt
      (2) INFORMATION FOR SEQ ID NO:107:
40
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 140 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
45
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                 (B) LOCATION: 1..140
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1665273
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:107:
      Met Ala Ala Thr Phe Ala Thr Pro Ser Thr Val Ile Gly Leu Gly Gly
                                          10
      Ser Ser Ile Thr Thr Lys Pro Phe Ser Ser Ser Phe Leu Lys Pro Thr
                                      25
55
      Leu Ser Ala Lys Asn Pro Leu Arg Leu Ala Gly Ala Ser Gly Gly Arg
                                  40
      Val Thr Cys Phe Glu Arg Asn Trp Leu Arg Arg Asp Leu Asn Val Val
                               55
      Gly Phe Gly Leu Ile Gly Trp Leu Ala Pro Ser Ser Ile Pro Ala Ile
60
                          70
                                           75
      Asn Gly Lys Ser Leu Thr Gly Leu Phe Phe Asp Ser Ile Gly Thr Glu
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Leu Ala His Phe Pro Thr Pro Pro Ala Leu Thr Ser Gln Phe Trp Leu
                                      105
     Trp Leu Val Thr Trp His Leu Gly Leu Phe Leu Cys Leu Thr Phe Gly
                                  120
 5
     Gln Ile Gly Phe Lys Gly Arg Thr Glu Asp Tyr Phe
                              135
      (2) INFORMATION FOR SEQ ID NO:108:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 522 base pairs
10
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
15
                (A) NAME/KEY: -
                (B) LOCATION: 1..522
                (D) OTHER INFORMATION: / Ceres Seq. ID 1713895
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:108:
      aaacttggat cttaacacag agaagaagca caatcggaag aaaggagaag acgaagatgt
                                                                             60
20
      cgttggtatg gctggaagcg atgttgcctc tcggaatcat cggtgggatg ctctgtatca
                                                                             120
      tgggcaattc tcagtactac atccacaaag cttatcatgg ccgtcctaag cacatcggcc
                                                                             180
      acgatgaatg ggatgttgct atggaaagac gcgacaagaa agtcgtcgag aaagctgcag
                                                                             240
      ctccttcctc atgattcgct ttatctcttt tgtgttcctc aggggcttaa ggtgaccact
                                                                             300
      tgtggtgaca aataaagtgc attccagaag aagaagaagc tgggggatct agtactttca
                                                                            360
25
      ttcccatttg attttccttg gacatattaa agctttcaga aatcagacct caataacatt
                                                                            420
      tggtttatca atatttctct attcgtgata ttttgtatgc ttttaagttg catgatacac
                                                                            480
      tgaactactc tcttgctgtg tgagtgaata aatgaatatc tg
      (2) INFORMATION FOR SEQ ID NO:109:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 83 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..83
                (D) OTHER INFORMATION: / Ceres Seq. ID 1713896
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:109:
40
      Thr Trp Ile Leu Thr Gln Arg Arg Ser Thr Ile Gly Arg Lys Glu Lys
                                          10
      Thr Lys Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile
                                       25
      Ile Gly Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His
45
                                   40
      Lys Ala Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp
      Val Ala Met Glu Arg Arg Asp Lys Lys Val Val Glu Lys Ala Ala Ala
50
      Pro Ser Ser
      (2) INFORMATION FOR SEQ ID NO:110:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 65 amino acids
55
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
           (ix) FEATURE:
60
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..65
```

(D) OTHER INFORMATION: / Ceres Seq. ID 1713897

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:110:
     Met Ser Leu Val Trp Leu Glu Ala Met Leu Pro Leu Gly Ile Ile Gly
                                          10
     Gly Met Leu Cys Ile Met Gly Asn Ser Gln Tyr Tyr Ile His Lys Ala
 5
                  20
                                      25
     Tyr His Gly Arg Pro Lys His Ile Gly His Asp Glu Trp Asp Val Ala
                                 40
     Met Glu Arg Arg Asp Lys Val Val Glu Lys Ala Ala Pro Ser
                              55
10
      Ser
      65
      (2) INFORMATION FOR SEQ ID NO:111:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 57 amino acids
15
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
20
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..57
                (D) OTHER INFORMATION: / Ceres Seq. ID 1713898
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:111:
      Met Leu Pro Leu Gly Ile Ile Gly Gly Met Leu Cys Ile Met Gly Asn
25
      1
                   5
                                          10
      Ser Gln Tyr Tyr Ile His Lys Ala Tyr His Gly Arg Pro Lys His Ile
                  20
                                      25
                                                           30
      Gly His Asp Glu Trp Asp Val Ala Met Glu Arg Arg Asp Lys Lys Val
                                  40
30
      Val Glu Lys Ala Ala Ala Pro Ser Ser
                              55
          50
      (2) INFORMATION FOR SEQ ID NO:112:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 745 base pairs
35
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
40
                (A) NAME/KEY: -
                (B) LOCATION: 1..745
                (D) OTHER INFORMATION: / Ceres Seq. ID 1923752
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:112:
                                                                              60
      acqcttaqqa tttaaqqact atcqttcttq qttqtqqatt ctcqcccacc qtcttqctta
45
      cgagtcacac accaccgtcc ttggatcgag tcttcaactt aagctctgat gctaattcac
                                                                             120
      atcaaagctg acttgctctg actttgtctc tcaaggattt gaggcgtgat tcaagaacaa
                                                                             180
      aacaaattaa agaacaactt ttctttattt atttagaaac tactcaaaac taaatctctc
                                                                             240
      taggtettta gaacaactet etaaageete teaatetett tettgagaet tatggaacaa
                                                                             300
      ctctccatag tctctctata tagaactgag ctacccctaa gtctatatct catgaagtct
                                                                             360
50
      ctctcqactc qcatqtttat ctcatqaqqt ctctctcqac tctaaqqaqa tctctcttqa
                                                                             420
      ctcacatctt gagetetete tttgateete ttggatgett tagetetett gtgttatett
                                                                             480
      cttcttcttc atagagatca tatatatact aggaccaaac acctcctttc cctattgtga
                                                                             540
      ataggaatac ttacaaatct tatccaactt gacattattt gtatttcttt ttttcttgat
                                                                             600
      ttcctttctt qaatttcttt tcctqtatct cttctaqcat cttctaqaaq qaqataactt
                                                                             660
55
      gctcctcaag taagttgtaa ttatccaaca cgcatgtgat ctttaatgat cgtggggatt
                                                                             720
      atttagtggc agagatgaat actct
      (2) INFORMATION FOR SEQ ID NO:113:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 35 amino acids
60
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
```

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643
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..35
 5
                (D) OTHER INFORMATION: / Ceres Seq. ID 1923753
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:113:
      Arg Leu Gly Phe Lys Asp Tyr Arg Ser Trp Leu Trp Ile Leu Ala His
                                          10
      Arg Leu Ala Tyr Glu Ser His Thr Thr Val Leu Gly Ser Ser Leu Gln
10
                  20
                                      25
      Leu Lys Leu
              35
      (2) INFORMATION FOR SEQ ID NO:114:
           (i) SEQUENCE CHARACTERISTICS:
15
                (A) LENGTH: 31 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
20
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..31
                (D) OTHER INFORMATION: / Ceres Seq. ID 1923754
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:114:
25
      Met Glu Gln Leu Ser Ile Val Ser Leu Tyr Arg Thr Glu Leu Pro Leu
                     5
                                          10
      Ser Leu Tyr Leu Met Lys Ser Leu Ser Thr Arg Met Phe Ile Ser
                  20
                                       25
      (2) INFORMATION FOR SEQ ID NO:115:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 36 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..36
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1923755
40
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:115:
      Met Arg Ser Leu Ser Thr Leu Arg Arg Ser Leu Leu Thr His Ile Leu
                                          10
      Ser Ser Leu Phe Asp Pro Leu Gly Cys Phe Ser Ser Leu Val Leu Ser
                  20
                                       25
45
      Ser Ser Ser Ser
              35
       (2) INFORMATION FOR SEQ ID NO:116:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 487 base pairs
50
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: DNA (genomic)
           (ix) FEATURE:
55
                 (A) NAME/KEY: -
                 (B) LOCATION: 1..487
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1976816
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:116:
      aaaccctagc aaaaacctct ctctctct caattcactt ctcctcctac aatggcttgg
                                                                               60
60
       cgcaacgcag gatctgctgc tcgttctttc gtctccgcca ccgcaagatc accatctctc
                                                                              120
       egttetecea ecaeggeget teetegeete egteeteete aateeteett aeetageegt
                                                                              180
```

egetteacet tetegteace ttecaqqaat etaggageee taggttgeac acagtegtte

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ttgcctctgt acagtgttgt ggctacttct caactgacat ctcaccttaa cgttaatttg
     cgagctttct gcgagctgtc taacggtact tgaagaggca aaagagtgta atttcactgc
                                                                             360
     gcatgcgtga atcagtactt tgaagggatg tgagatcgaa attgtgggat tacctcagca
                                                                             420
     aaaacttgct taatgtgttg cctgttttag aagttttagt ttcacaatgt gtggatttat
                                                                             480
 5
     tagttgt
      (2) INFORMATION FOR SEQ ID NO:117:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 70 amino acids
                (B) TYPE: amino acid
10
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
15
                (B) LOCATION: 1..70
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976817
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:117:
      Asn Pro Ser Lys Asn Leu Ser Leu Ser Leu Asn Ser Leu Leu Leu
                                          10
                      5
20
      Gln Trp Leu Gly Ala Thr Gln Asp Leu Leu Leu Val Leu Ser Ser Pro
                  20
                                     25
      Pro Pro Gln Asp His His Leu Ser Val Leu Pro Pro Arg Arg Phe Leu
                                  40
      Ala Ser Val Leu Leu Asn Pro Pro Tyr Leu Ala Val Ala Ser Pro Ser
25
                              55
                                                   60
      Arg His Leu Pro Gly Ile
                          70
      (2) INFORMATION FOR SEQ ID NO:118:
           (i) SEQUENCE CHARACTERISTICS:
30
                (A) LENGTH: 57 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
35
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..57
                (D) OTHER INFORMATION: / Ceres Seq. ID 1976818
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:118:
40
      Thr Leu Ala Lys Thr Ser Leu Ser Leu Ser Ile His Phe Ser Ser Tyr
                                          10
      Asn Gly Leu Ala Gln Arg Arg Ile Cys Cys Ser Phe Phe Arg Leu Arg
                                       25
      His Arg Lys Ile Thr Ile Ser Pro Phe Ser His His Gly Ala Ser Ser
45
                                  40
              35
      Pro Pro Ser Ser Ser Ile Leu Leu Thr
      (2) INFORMATION FOR SEQ ID NO:119:
           (i) SEQUENCE CHARACTERISTICS:
50
                 (A) LENGTH: 93 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
55
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..93
                 (D) OTHER INFORMATION: / Ceres Seq. ID 1976819
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:119:
60
      Met Ala Trp Arg Asn Ala Gly Ser Ala Ala Arg Ser Phe Val Ser Ala
                                           10
      Thr Ala Arg Ser Pro Ser Leu Arg Ser Pro Thr Thr Ala Leu Pro Arg
```

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20 25 30

Leu Arg Pro Pro Gln Ser Ser Leu Pro Ser Arg Arg Phe Thr Phe Ser 35 40 45

Ser Pro Ser Arg Asn Leu Gly Ala Leu Gly Cys Thr Gln Ser Phe Leu 50 55 60

Pro Leu Tyr Ser Val Val Ala Thr Ser Gln Leu Thr Ser His Leu Asn

65 70 75 80
Val Asn Leu Arg Ala Phe Cys Glu Leu Ser Asn Gly Thr

- 85
  10 (2) INFORMATION FOR SEQ ID NO:120:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 566 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
- 15 (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (ix) FEATURE:

5

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- (A) NAME/KEY: -
- (B) LOCATION: 1..566
- 20 (D) OTHER INFORMATION: / Ceres Seq. ID 2025128
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:120:

atgaatette ttegettegg eteegeetet tattetegag aaageteate ttgettetea 60 catctaaaca aatatctcat ctcttctatt ctactttatc acttcatctc cgtacgctct 120 totttottog ttgottotto ttttgattta taaattggat ccaaaacatc ataggototo 180 gtcatcctta tatacaagca ttgttacttt gtgatataaa gaacgaagaa ttgtacgaag 240 gcaacgagca agatgagtag caaggcgaat gaggtttaaa agagtagcaa ggcgaatgaa 300 gttgaaagga ggaagatgaa tagcacggcc aatgacgatg aaaggatgaa taggatcatc 360 agagatggcc gtcgtcgtta cgtcacctgg agccgcagcc gcaacccctc ggcagctcat 420 atcttttggt caaaacccta atattttcta gggttgggct attagttttt gttttgggct 480

540

- 30 tttttgaatt gtattttggg ctttttatta atttgtagtg ggtaatttgt ttggattagg atttttggta gtgtaatttt tttttt
  - (2) INFORMATION FOR SEQ ID NO:121:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 50 amino acids
      - (B) TYPE: amino acid
      - (C) STRANDEDNESS:
      - (D) TOPOLOGY: linear
    - (ii) MOLECULE TYPE: peptide
    - (ix) FEATURE:
      - (A) NAME/KEY: peptide
        - (B) LOCATION: 1..50
        - (D) OTHER INFORMATION: / Ceres Seq. ID 2025129
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:121:

Met Asn Leu Leu Arg Phe Gly Ser Ala Ser Tyr Ser Arg Glu Ser Ser 45 1 5 10 15

Ser Cys Phe Ser His Leu Asn Lys Tyr Leu Ile Ser Ser Ile Leu Leu 20 25 30

Tyr His Phe Ile Ser Val Arg Ser Ser Phe Phe Val Ala Ser Ser Phe 35 40 45

- 50 Asp Leu 50
  - (2) INFORMATION FOR SEQ ID NO:122:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 47 amino acids
    - (B) TYPE: amino acid
      - (C) STRANDEDNESS:
      - (D) TOPOLOGY: linear
    - (ii) MOLECULE TYPE: peptide
    - (ix) FEATURE:
  - (A) NAME/KEY: peptide
    - (B) LOCATION: 1..47
    - (D) OTHER INFORMATION: / Ceres Seq. ID 2025130

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:122:
      Glu Ser Ser Ser Leu Arg Leu Arg Leu Leu Phe Ser Arg Lys Leu Ile
                                          10
      Leu Leu Thr Ser Lys Gln Ile Ser His Leu Phe Tyr Ser Thr Leu
               20
 5
                                   25
      Ser Leu His Leu Arg Thr Leu Phe Phe Leu Arg Cys Phe Phe Phe
                                  40
      (2) INFORMATION FOR SEQ ID NO:123:
           (i) SEQUENCE CHARACTERISTICS:
10
                (A) LENGTH: 41 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
15
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..41
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025131
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:123:
20
      Met Asn Ser Thr Ala Asn Asp Asp Glu Arg Met Asn Arg Ile Ile Arg
                                           10
      Asp Gly Arg Arg Arg Tyr Val Thr Trp Ser Arg Ser Arg Asn Pro Ser
                                    25
                  20
      Ala Ala His Ile Phe Trp Ser Lys Pro
25
              35
      (2) INFORMATION FOR SEQ ID NO:124:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 545 base pairs
                 (B) TYPE: nucleic acid
30
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                 (A) NAME/KEY: -
35
                 (B) LOCATION: 1..545
                 (D) OTHER INFORMATION: / Ceres Seq. ID 2025402
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:124:
      ttttttttt ttttgcgaaa ttgaaatgat ctcacataaa aactcaattg ataattttac
      aaaggagaaa aatttgtcaa tcaaagaaga agagaaagat tgaactttta acctcacttg
                                                                               120
40
      gtcataactt caacgccatt ctttccagcg ataatgacag aagtagccat tccgagaaac
                                                                               180
      agaccatgct ccaccactcc ttgaaacttc ccaatctctt tagccgcagc gaatccatcc
                                                                               240
      ttcaaaggag tcttaaaata caaatcaata atgtaattac tgttatcagt cacataaggc
                                                                               300
      ttgccatcac catcaactct aagctttgat tcacatccaa attccttgaa gaggtcttgc
                                                                               360
      aatctaatca aattaaagtt ccagcagaat tgaacaactt ccaccggcat agctaatcca cttccaccga gtcctgtaac gagtttggta tcatcagcca caacaataaa cttgtcagcc
                                                                              420
45
                                                                              480
                                                                              540
      acagetteca ceattaceta gaacateeta ategaaaaat tateeaatee aateeattae
      ttact
      (2) INFORMATION FOR SEQ ID NO:125:
            (i) SEQUENCE CHARACTERISTICS:
50
                 (A) LENGTH: 47 amino acids
                 (B) TYPE: amino acid
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: peptide
55
           (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..47
                 (D) OTHER INFORMATION: / Ceres Seq. ID 2025403
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:125:
60
      Met Leu His His Ser Leu Lys Leu Pro Asn Leu Phe Ser Arg Ser Glu
                                           10
       Ser Ile Leu Gln Arg Ser Leu Lys Ile Gln Ile Asn Asn Val Ile Thr
```

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25
                                                          30
                  20
      Val Ile Ser His Ile Arg Leu Ala Ile Thr Ile Asn Ser Lys Leu
              3.5
                                  40
      (2) INFORMATION FOR SEQ ID NO:126:
 5
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 33 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
10
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..33
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025404
15
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:126:
      Phe Phe Phe Cys Glu Ile Glu Met Ile Ser His Lys Asn Ser Ile
                                          10
      Asp Asn Phe Thr Lys Glu Lys Asn Leu Ser Ile Lys Glu Glu Glu Lys
                                      25
                  20
20
      Asp
      (2) INFORMATION FOR SEQ ID NO:127:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1954 base pairs
25
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
30
                (A) NAME/KEY: -
                (B) LOCATION: 1..1954
                (D) OTHER INFORMATION: / Ceres Seq. ID 2025479
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:127:
      atggaacctt atataaccag aagagataca gttaatcaag tgttacatgc agaagaagaa
                                                                             60
35
      gaagaataca atgattgtga attggatcaa ctcttccttg tgcattctga tattagatct
                                                                             120
      gttcttctcc agattgatga gcttgttgtt gaagccacaa agcgtaagac cgtgagcaag
                                                                             180
      cacgggttaa tagaagttga atctttcaga actgtgttgt ctgatatgct ttcctcacta
                                                                             240
      aagccatggt ttcctcgatt gcaagaagcg atgtctgatt ttcagttgct tccagaggat
                                                                             300
                                                                             360
      caggaagaac agtctttgat gagcacaaac gaagaagaag atttgtttga tgttgagagt
40
                                                                             420
      cctqaqccta ctcaatttqa gcccttaqtc tctccttctc ctcttqtqca ttqqcqtgqt
      gatcataacg cagacaaggg tagacaactc tttcttttga caccactacc cttaggaaaa
                                                                             480
                                                                             540
      tctgaattcc taaaacacca gaatgcatct aagcttacag ctaaaaggat ttttccagac
                                                                             600
      actgttgcaa atgaaccgct tgaagcttct aaagaaacaa gtgatgatgt tttaggaggt
                                                                             660
      gagtcgttga aaactgcagg acttggtaaa tctttagtcc atgccatgga tttctctgaa
45
                                                                             720
      aatttggttg agtataagcc atgttcttca ccagttctca ggagaaagat tctgtctgag
                                                                             780
      cttctaatga caccttgctt gaagctgtcg cctccgaaat cgtgtacaat gtttaaaccg
                                                                             840
      gttcctgaat cttctcaact gggaaagcaa ggggcttgta agtctacctg ctctgaattg
                                                                             900
      ggatcatctg gtattgaaaa aacagacaat ctctgctcga agtatcctga gctcttggga
                                                                             960
      atacaacatg ctccgataac tagaaaaacg gatcttgaat cctcgccggt ttggtggttt
50
                                                                            1020
      tctcctccta aaacatgtgt tttaatggag cctgtgaatg agaagaaacc aattgatgaa
      actggtggat cttttgatgt tccaaacatt ataccagaag caaaacacac tacagaggga
                                                                            1080
      agcatgtcaa tggtggttga gagtactcca ttgtttaaag aaccagagag cataatgacg
                                                                            1140
      agaaacagaa caaaagcagg tgagagtact ctgaagaaag agctatggac aagatttgaa
                                                                            1200
      qaaqcaacqa ttcatqacaq tcqqttcaac tcqatqacaa caacaacaac aqtqaqaqqa
                                                                            1260
55
      aataataaga aatgtttcat ggaaatgtta gaagaagtga gtggcaatga ggaagatcat
                                                                            1320
      gagttaagtg gcgagtttgt ggctctaaaa ccttatcaag tactcatatc ggaactgagc
                                                                            1380
      ataacaaacg cttttcattg tctgtctact cttattagct gctacataac tcaaagtaac
                                                                            1440
      aaacatacac atataacata tagaaagatc gaaaacaaaa gatcgggatg agtatggctc
                                                                            1500
      caaagacctc aactacactt gctttattcc ttgtgaccaa tattctcttc ctcaacctca
60
      ttaccctgag ttgcgcagac aatacttgcc caagagacgt tctcaaactt tcgacatgct
                                                                            1620
      cgaatgttct caacctcatc aacttgaagc tcggggcacc agctatgagg ccttgttgct
                                                                            1680
                                                                           1740
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ctattetett tggtetaatt gatetegatg ttgeggtttg cetttgeace gegeteaage

648

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gtggaggtac cetteetgat ggatteegtt geecaacata getacaatat atteatattt tetatetatg teettgaatt ggtttTgtet aacateeaaa tteaaaceta aatteatgta 1920 taageaaata acaagtaaaa ttaageatt teettgaatt teettga
         taagcaaata acaagtaaaa ttaagacttt tatg
 5
         (2) INFORMATION FOR SEQ ID NO:128:
                  (i) SEQUENCE CHARACTERISTICS:
                           (A) LENGTH: 496 amino acids
                          (B) TYPE: amino acid
                          (C) STRANDEDNESS:
10
                          (D) TOPOLOGY: linear
                (ii) MOLECULE TYPE: peptide
                (ix) FEATURE:
                          (A) NAME/KEY: peptide
                          (B) LOCATION: 1..496
15
                          (D) OTHER INFORMATION: / Ceres Seq. ID 2025480
                 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:128:
         Met Glu Pro Tyr Ile Thr Arg Arg Asp Thr Val Asn Gln Val Leu His
                                                                     10
          Ala Glu Glu Glu Glu Tyr Asn Asp Cys Glu Leu Asp Gln Leu Phe
20
                                                                25
          Leu Val His Ser Asp Ile Arq Ser Val Leu Leu Gln Ile Asp Glu Leu
                                                         40
          Val Val Glu Ala Thr Lys Arg Lys Thr Val Ser Lys His Gly Leu Ile
                                                  55
25
          Glu Val Glu Ser Phe Arg Thr Val Leu Ser Asp Met Leu Ser Ser Leu
                                            70
                                                                              75
          Lys Pro Trp Phe Pro Arg Leu Gln Glu Ala Met Ser Asp Phe Gln Leu
                                                                       90
          Leu Pro Glu Asp Gln Glu Glu Gln Ser Leu Met Ser Thr Asn Glu Glu
                                                               105
30
                              100
          Glu Asp Leu Phe Asp Val Glu Ser Pro Glu Pro Thr Gln Phe Glu Pro
                      115
                                                        120
                                                                                          125
          Leu Val Ser Pro Ser Pro Leu Val His Trp Arg Gly Asp His Asn Ala
                                                                                   140^{-}
                                                 135
35
          Asp Lys Gly Arg Gln Leu Phe Leu Leu Thr Pro Leu Pro Leu Gly Lys
                                          150 155
          Ser Glu Phe Leu Lys His Gln Asn Ala Ser Lys Leu Thr Ala Lys Arg
                                    165 170 175
          Ile Phe Pro Asp Thr Val Ala Asn Glu Pro Leu Glu Ala Ser Lys Glu
40
                             180
          Thr Ser Asp Asp Val Leu Gly Gly Glu Ser Leu Lys Thr Ala Gly Leu
                                                        200
          Gly Lys Ser Leu Val His Ala Met Asp Phe Ser Glu Asn Leu Val Glu
                                                 215
                                                                                   220
45
          Tyr Lys Pro Cys Ser Ser Pro Val Leu Arg Arg Lys Ile Leu Ser Glu
                                           230
                                                                            235 240
          Leu Leu Met Thr Pro Cys Leu Lys Leu Ser Pro Pro Lys Ser Cys Thr
                                    245
                                                                      250 255
          Met Phe Lys Pro Val Pro Glu Ser Ser Gln Leu Gly Lys Gln Gly Ala
50
                                                               265
          Cys Lys Ser Thr Cys Ser Glu Leu Gly Ser Ser Gly Ile Glu Lys Thr
                                                         280
          Asp Asn Leu Cys Ser Lys Tyr Pro Glu Leu Leu Gly Ile Gln His Ala
                                                  295
55
          Pro Ile Thr Arg Lys Thr Asp Leu Glu Ser Ser Pro Val Trp Trp Phe
                                                                             315
                                           310
          Ser Pro Pro Lys Thr Cys Val Leu Met Glu Pro Val Asn Glu Lys Lys
                                                                       330
           Pro Ile Asp Glu Thr Gly Gly Ser Phe Asp Val Pro Asn Ile Ile Pro
 60
                             340 345 350
           Glu Ala Lys His Thr Thr Glu Gly Ser Met Ser Met Val Val Glu Ser
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Thr Pro Leu Phe Lys Glu Pro Glu Ser Ile Met Thr Arg Asn Arg Thr
                         375
    Lys Ala Gly Glu Ser Thr Leu Lys Lys Glu Leu Trp Thr Arg Phe Glu
                      390
                                      395
    Glu Ala Thr Ile His Asp Ser Arg Phe Asn Ser Met Thr Thr Thr
                      410 415
     Thr Val Arg Gly Asn Asn Lys Lys Cys Phe Met Glu Met Leu Glu Glu
                   425 430
     Val Ser Gly Asn Glu Glu Asp His Glu Leu Ser Gly Glu Phe Val Ala
10
           435
                            440 445
     Leu Lys Pro Tyr Gln Val Leu Ile Ser Glu Leu Ser Ile Thr Asn Ala
                         455 460
     Phe His Cys Leu Ser Thr Leu Ile Ser Cys Tyr Ile Thr Gln Ser Asn
        470 475
15
     Lys His Thr His Ile Thr Tyr Arg Lys Ile Glu Asn Lys Arg Ser Gly
                          490
                  485
     (2) INFORMATION FOR SEQ ID NO:129:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 421 amino acids
20
             (B) TYPE: amino acid
             (C) STRANDEDNESS:
             (D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: peptide
        (ix) FEATURE:
25
             (A) NAME/KEY: peptide
             (B) LOCATION: 1..421
             (D) OTHER INFORMATION: / Ceres Seq. ID 2025481
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:129:
     Met Leu Ser Ser Leu Lys Pro Trp Phe Pro Arg Leu Gln Glu Ala Met
30
                                   10
     Ser Asp Phe Gln Leu Leu Pro Glu Asp Gln Glu Glu Gln Ser Leu Met
         20 25
     Ser Thr Asn Glu Glu Glu Asp Leu Phe Asp Val Glu Ser Pro Glu Pro
                            40
35
     Thr Gln Phe Glu Pro Leu Val Ser Pro Ser Pro Leu Val His Trp Arg
                      55
                                       60
     Gly Asp His Asn Ala Asp Lys Gly Arg Gln Leu Phe Leu Leu Thr Pro
                     70
                                      75 80
     Leu Pro Leu Gly Lys Ser Glu Phe Leu Lys His Gln Asn Ala Ser Lys
40
                  85 90
     Leu Thr Ala Lys Arg Ile Phe Pro Asp Thr Val Ala Asn Glu Pro Leu
                                105
     Glu Ala Ser Lys Glu Thr Ser Asp Asp Val Leu Gly Glv Glu Ser Leu
                            120 125
45
     Lys Thr Ala Gly Leu Gly Lys Ser Leu Val His Ala Met Asp Phe Ser
                        135
     Glu Asn Leu Val Glu Tyr Lys Pro Cys Ser Ser Pro Val Leu Arg Arg
                     150
     Lys Ile Leu Ser Glu Leu Leu Met Thr Pro Cys Leu Lys Leu Ser Pro
50
                  165
                                   170
     Pro Lys Ser Cys Thr Met Phe Lys Pro Val Pro Glu Ser Ser Gln Leu
                                185
     Gly Lys Gln Gly Ala Cys Lys Ser Thr Cys Ser Glu Leu Gly Ser Ser
           195 200
55
     Gly Ile Glu Lys Thr Asp Asn Leu Cys Ser Lys Tyr Pro Glu Leu Leu
                         215
                                          220
     Gly Ile Gln His Ala Pro Ile Thr Arg Lys Thr Asp Leu Glu Ser Ser
                     230
                                       235
     Pro Val Trp Trp Phe Ser Pro Pro Lys Thr Cys Val Leu Met Glu Pro
60
               245 250
     Val Asn Glu Lys Lys Pro Ile Asp Glu Thr Gly Gly Ser Phe Asp Val
               260
                                265
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650 Pro Asn Ile Ile Pro Glu Ala Lys His Thr Thr Glu Gly Ser Met Ser 280 285 Met Val Val Glu Ser Thr Pro Leu Phe Lys Glu Pro Glu Ser Ile Met 295 300 5 Thr Arg Asn Arg Thr Lys Ala Gly Glu Ser Thr Leu Lys Lys Glu Leu 310 315 Trp Thr Arg Phe Glu Glu Ala Thr Ile His Asp Ser Arg Phe Asn Ser 330 335 Met Thr Thr Thr Thr Val Arg Gly Asn Asn Lys Lys Cys Phe Met 10 340 345 350 Glu Met Leu Glu Glu Val Ser Gly Asn Glu Glu Asp His Glu Leu Ser 360 365 Gly Glu Phe Val Ala Leu Lys Pro Tyr Gln Val Leu Ile Ser Glu Leu 370 375 380 15 Ser Ile Thr Asn Ala Phe His Cys Leu Ser Thr Leu Ile Ser Cys Tyr 390 395 Ile Thr Gln Ser Asn Lys His Thr His Ile Thr Tyr Arg Lys Ile Glu 405 410 415 Asn Lys Arg Ser Gly 20 420 (2) INFORMATION FOR SEQ ID NO:130: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 406 amino acids (B) TYPE: amino acid
(C) STRANDEDNESS: 25 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 30 (B) LOCATION: 1..406 (D) OTHER INFORMATION: / Ceres Seq. ID 2025482 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:130: Met Ser Asp Phe Gln Leu Leu Pro Glu Asp Gln Glu Glu Gln Ser Leu 10 35 Met Ser Thr Asn Glu Glu Glu Asp Leu Phe Asp Val Glu Ser Pro Glu 20 25 Pro Thr Gln Phe Glu Pro Leu Val Ser Pro Ser Pro Leu Val His Trp 40 Arg Gly Asp His Asn Ala Asp Lys Gly Arg Gln Leu Phe Leu Leu Thr 40 55 60 Pro Leu Pro Leu Gly Lys Ser Glu Phe Leu Lys His Gln Asn Ala Ser 70 75 80 Lys Leu Thr Ala Lys Arg Ile Phe Pro Asp Thr Val Ala Asn Glu Pro 90 95 45 Leu Glu Ala Ser Lys Glu Thr Ser Asp Asp Val Leu Gly Gly Glu Ser 105 Leu Lys Thr Ala Gly Leu Gly Lys Ser Leu Val His Ala Met Asp Phe 120 Ser Glu Asn Leu Val Glu Tyr Lys Pro Cys Ser Ser Pro Val Leu Arg 50 135 Arg Lys Ile Leu Ser Glu Leu Leu Met Thr Pro Cys Leu Lys Leu Ser 150 155 160 Pro Pro Lys Ser Cys Thr Met Phe Lys Pro Val Pro Glu Ser Ser Gln 165 170 55 Leu Gly Lys Gln Gly Ala Cys Lys Ser Thr Cys Ser Glu Leu Gly Ser 180 185 Ser Gly Ile Glu Lys Thr Asp Asn Leu Cys Ser Lys Tyr Pro Glu Leu 200 205 Leu Gly Ile Gln His Ala Pro Ile Thr Arg Lys Thr Asp Leu Glu Ser 60 210 215 220 Ser Pro Val Trp Trp Phe Ser Pro Pro Lys Thr Cys Val Leu Met Glu

230

225

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651
     Pro Val Asn Glu Lys Lys Pro Ile Asp Glu Thr Gly Gly Ser Phe Asp
                      245
                                          250
     Val Pro Asn Ile Ile Pro Glu Ala Lys His Thr Thr Glu Gly Ser Met
                                      265
5
     Ser Met Val Val Glu Ser Thr Pro Leu Phe Lys Glu Pro Glu Ser Ile
                                  280
     Met Thr Arg Asn Arg Thr Lys Ala Gly Glu Ser Thr Leu Lys Lys Glu
                              295
                                                  300
     Leu Trp Thr Arg Phe Glu Glu Ala Thr Ile His Asp Ser Arg Phe Asn
10
                          310
                                              315
     Ser Met Thr Thr Thr Thr Val Arg Gly Asn Asn Lys Lys Cys Phe
                      325
                                          330
     Met Glu Met Leu Glu Glu Val Ser Gly Asn Glu Glu Asp His Glu Leu
                  340
                                      345
15
     Ser Gly Glu Phe Val Ala Leu Lys Pro Tyr Gln Val Leu Ile Ser Glu
                                  360
     Leu Ser Ile Thr Asn Ala Phe His Cys Leu Ser Thr Leu Ile Ser Cys
                              375
                                                  380
     Tyr Ile Thr Gln Ser Asn Lys His Thr His Ile Thr Tyr Arg Lys Ile
20
                         390
                                              395
     Glu Asn Lys Arg Ser Gly
                      405
      (2) INFORMATION FOR SEQ ID NO:131:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 609 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
30
          (ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..609
                (D) OTHER INFORMATION: / Ceres Seq. ID 2032963
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:131:
35
                                                                             60
     tttgtgtggt aattaactgg tgagagagtt caacatccaa tctttctctc tctctcta
      tottttatot otoacactot taaaatgott ottotatoto ogatototgo ototottooa
                                                                            120
                                                                            180
      ccgtcgtttc accggggtaa tttgatccgg cgatcaatta agccgttagg tagggttgtt
      gcaaaggcca aggacaacac cgacactggt ggatttctgg agacggcagc tatagccggt
                                                                            240
      agtttagtgt cgacgccggt tattggatgg tcactctaca ctttgaagac cacgggatgt
                                                                            300
40
      ggcttacctc ccggaccggc cggtttaatc ggagcgttgg aaggtgtgag ctacttggtg
                                                                            360
      gtggttggca tcgtgggctg gtctttgtac actaaaacaa aaactgggtc aggtctgcca
                                                                            420
      aatgggccat ttggtttgtt gggtgcggtc gagggtttat cgtatctgtc ggttctagcc
                                                                            480
      attettgtgg tgtttggtat teagttettg gataatgggt eggtteeagg teeactteet
                                                                            540
      agtgaccagt gttttggtta accacgtgta atgtcgtata aataaagtgt caaattaata
                                                                            600
45
      attgctttc
      (2) INFORMATION FOR SEQ ID NO:132:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 158 amino acids
                (B) TYPE: amino acid
50
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
55
                (B) LOCATION: 1..158
                (D) OTHER INFORMATION: / Ceres Seq. ID 2032964
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:132:
      Met Leu Leu Ser Pro Ile Ser Ala Ser Leu Pro Pro Ser Phe His
                      5
                                          10
60
      Arg Gly Asn Leu Ile Arg Arg Ser Ile Lys Pro Leu Gly Arg Val Val
                                      25
      Ala Lys Ala Lys Asp Asn Thr Asp Thr Gly Gly Phe Leu Glu Thr Ala
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652
                                  40
      Ala Ile Ala Gly Ser Leu Val Ser Thr Pro Val Ile Gly Trp Ser Leu
                              55
                                                  60
      Tyr Thr Leu Lys Thr Thr Gly Cys Gly Leu Pro Pro Gly Pro Ala Gly
 5
                          70
                                              75
      Leu Ile Gly Ala Leu Glu Gly Val Ser Tyr Leu Val Val Val Gly Ile
                     8.5
                                          90
      Val Gly Trp Ser Leu Tyr Thr Lys Thr Lys Thr Gly Ser Gly Leu Pro
                  100
                                     105
                                                          110
10
     Asn Gly Pro Phe Gly Leu Leu Gly Ala Val Glu Gly Leu Ser Tyr Leu
                                  120
      Ser Val Leu Ala Ile Leu Val Val Phe Gly Ile Gln Phe Leu Asp Asn
                          135
                                                  140
      Gly Ser Val Pro Gly Pro Leu Pro Ser Asp Gln Cys Phe Gly
15
                         150 ..
      (2) INFORMATION FOR SEQ ID NO:133:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 330 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
25
                (B) LOCATION: 1..330
                (D) OTHER INFORMATION: / Ceres Seq. ID 2033706
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:133:
      atgggccaca gagtgacgag agagagagtg aaaatggaga agtactttgg aaatgcgtac
                                                                             60
      aggggagate caggagtgee acatgeagat geagategtt tegtgaatat atggattggt
30
      totgctqctt totccqttct tacctqqqtt aatccttaca tqtqqcaqct ctctaaccaq
                                                                           180
      ttcaattacc atgacaagtg gatgctgttt gagcagtacc actggaaaaa agcaagggca
                                                                             240
      aaqaaqcaac cttatqaatt caaqtqqaat aaqataccca aaqaaqtcaq qqactcqtac
                                                                             300
      tattacaact ggccagtcta cttcccatag
      (2) INFORMATION FOR SEQ ID NO:134:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 109 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..109
                (D) OTHER INFORMATION: / Ceres Seq. ID 2033707
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:134:
      Met Gly His Arg Val Thr Arg Glu Arg Val Lys Met Glu Lys Tyr Phe
                      5
                                          10
                                                               1.5
      Gly Asn Ala Tyr Arg Gly Asp Pro Gly Val Pro His Ala Asp Ala Asp
                  20
                                      25
50
      Arg Phe Val Asn Ile Trp Ile Gly Ser Ala Ala Phe Ser Val Leu Thr
                                  40
      Trp Val Asn Pro Tyr Met Trp Gln Leu Ser Asn Gln Phe Asn Tyr His
          50
                              55
      Asp Lys Trp Met Leu Phe Glu Gln Tyr His Trp Lys Lys Ala Arg Ala
55
                                              75
                          70
      Lys Lys Gln Pro Tyr Glu Phe Lys Trp Asn Lys Ile Pro Lys Glu Val
                      85
                                          90
      Arg Asp Ser Tyr Tyr Tyr Asn Trp Pro Val Tyr Phe Pro
                  100
60
      (2) INFORMATION FOR SEQ ID NO:135:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 98 amino acids

653 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 5 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..98 (D) OTHER INFORMATION: / Ceres Seq. ID 2033708 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:135: 10 Met Glu Lys Tyr Phe Gly Asn Ala Tyr Arg Gly Asp Pro Gly Val Pro 10 His Ala Asp Ala Asp Arg Phe Val Asn Ile Trp Ile Gly Ser Ala Ala 20 25 Phe Ser Val Leu Thr Trp Val Asn Pro Tyr Met Trp Gln Leu Ser Asn 15 4.0 4.5 Gln Phe Asn Tyr His Asp Lys Trp Met Leu Phe Glu Gln Tyr His Trp 55 Lys Lys Ala Arg Ala Lys Lys Gln Pro Tyr Glu Phe Lys Trp Asn Lys 70 75 80 20 Ile Pro Lys Glu Val Arg Asp Ser Tyr Tyr Tyr Asn Trp Pro Val Tyr 90 Phe Pro (2) INFORMATION FOR SEQ ID NO:136: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 56 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: Linear 30 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..56 (D) OTHER INFORMATION: / Ceres Seq. ID 2033709 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:136: Met Trp Gln Leu Ser Asn Gln Phe Asn Tyr His Asp Lys Trp Met Leu 5 10 Phe Glu Gln Tyr His Trp Lys Lys Ala Arg Ala Lys Lys Gln Pro Tyr 25 2.0 30 40 Glu Phe Lys Trp Asn Lys Ile Pro Lys Glu Val Arg Asp Ser Tyr Tyr 40 Tyr Asn Trp Pro Val Tyr Phe Pro 50 (2) INFORMATION FOR SEQ ID NO:137: 45 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 418 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 50 (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -(B) LOCATION: 1..418 (D) OTHER INFORMATION: / Ceres Seq. ID 2043118 55 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:137: ataacaatat caaaagataa aaacagagtt tgcttttata agatcaaaga aggtaaaaaa agaaatggca aaatggagtg cgatcgtatt gataatgatg gtgatgattg ttgcggtcac 120 aatagaagca caagaagaaa gtggtgggac tatttgtttt cqccaatgtt ctcagccttg 180 tegeagtgae gaeggtagtt getatgaaaa etgtaaaate gaatgeggtg gteecaagee 240 60 tecettatet egtetaegaa geteaeatge atgagatgge ateateaatg gaagttegeg 300 gaaggatagg atgataaaaa gagatttgta gcggattttt gatcatctct actgttttta 360

acttcacgtt ttatattata agagtttcat taaaaagatc aataaaacga gaaaatgt

PCT/US00/00466

WO 00/40695 654 (2) INFORMATION FOR SEQ ID NO:138: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 69 amino acids (B) TYPE: amino acid 5 (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide 10 (B) LOCATION: 1..69 (D) OTHER INFORMATION: / Ceres Seq. ID 2043119 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:138: Met Ala Lys Trp Ser Ala Ile Val Leu Ile Met Met Val Met Ile Val 5 10 15 Ala Val Thr Ile Glu Ala Gln Glu Glu Ser Gly Gly Thr Ile Cys Phe 20 25 Arg Gln Cys Ser Gln Pro Cys Arg Ser Asp Asp Gly Ser Cys Tyr Glu 40 Asn Cys Lys Ile Glu Cys Gly Gly Pro Lys Pro Pro Leu Ser Arg Leu 20 50 55 60 Arg Ser Ser His Ala (2) INFORMATION FOR SEQ ID NO:139: (i) SEQUENCE CHARACTERISTICS: 25 (A) LENGTH: 62 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 30 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..62 (D) OTHER INFORMATION: / Ceres Seq. ID 2043120 (xi) SEQUENCL DESCRIPTION: SEQ ID NO:139: 35 Ile Thr Ile Ser Lys Asp Lys Asn Arg Val Cys Phe Tyr Lys Ile Lys 10 Glu Gly Lys Lys Arg Asn Gly Lys Met Glu Cys Asp Arg Ile Asp Asn 20 25 Asp Gly Asp Asp Cys Cys Gly His Asn Arg Ser Thr Arg Arg Lys Trp 40 40 Trp Asp Tyr Leu Phe Ser Pro Met Phe Ser Ala Leu Ser Gln 50 55 (2) INFORMATION FOR SEQ ID NO:140: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 59 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 50 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..59 (D) OTHER INFORMATION: / Ceres Seq. ID 2043121 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:140: 55 Met Met Val Met Ile Val Ala Val Thr Ile Glu Ala Gln Glu Ser

Gly Gly Thr Ile Cys Phe Arg Gln Cys Ser Gln Pro Cys Arg Ser Asp 25 Asp Gly Ser Cys Tyr Glu Asn Cys Lys Ile Glu Cys Gly Gly Pro Lys 60 40 Pro Pro Leu Ser Arg Leu Arg Ser Ser His Ala 55

PCT/US00/00466 WO 00/40695 655 (2) INFORMATION FOR SEQ ID NO:141: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 838 base pairs (B) TYPE: nucleic acid 5 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (ix) FEATURE: (A) NAME/KEY: -10 (B) LOCATION: 1..838 (D) OTHER INFORMATION: / Ceres Seq. ID 2047214 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:141: acacgaaacg attocgcttc tocgcttoct toataaatat otoactoato gtototagat 60 tgttctttca cacgcaattt tcatttctct cttaaccgta aacgaatctc tcttctcaaa 120 15 gttttatttt ctctctgcga tggctcaaga agatgttact gctgttgcta ctaacggtgc 180 tggtccggtg gagacgcatc ttgtcttcac tgagttcaag cagatgttgc tcgttgaagc 240 300 tcaaaaggtc ggtgacgctg ttactttcta caaatctgct tttggtgcga tcgagtctgg teattetett taccetaage gtaagettga ceaagagett ceteatgtte tetettetga 360 gettaatete getggetett cettegttgt ttgegaegtt teetetee etggttttte 420 20 tactgcgaaa tcggaaggtt cgggagtgac ttttcttctc ggaactaagg atgctgaagc 480 cgccgttgcg aaagctgttg acgccggagc tgtgaaagtg gaggttacgg aggcagaagt 540 tgaactggga ttcaaaggaa aagttacgga teettttggt gteacttgga tettegegga 600 gaagaagacc gtgatcaccg acgagaacaa agaggtttag aatctgtcgt cggatctaat tcttcqtcqq tttctqaaca aaaaaaacta tctattatcc tagqtttatq cttttatttt 720 25 qtcttttqtq qaatccqqta atcaqtaaac cqqatcqaqt aqqctaatqa ctttcqqatt ctaatttcac cccttttgac aaactctaat cttggtgaaa ttgcatatta atctcggg (2) INFORMATION FOR SEQ ID NO:142: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 166 amino acids 30 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FLATURE: 35 (A) NAME/KEY: peptide (B) LOCATION: 1..166 (D) OTHER INFORMATION: / Ceres Seq. ID 2047215 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:142: Met Ala Gln Glu Asp Val Thr Ala Val Ala Thr Asn Gly Ala Gly Pro 40 10 Val Glu Thr His Leu Val Phe Thr Glu Phe Lys Gln Met Leu Leu Val 20 25 Glu Ala Gln Lys Val Gly Asp Ala Val Thr Phe Tyr Lys Ser Ala Phe 4.0 45 Gly Ala Ile Glu Ser Gly His Ser Leu Tyr Pro Lys Arg Lys Leu Asp 55 60 Gln Glu Leu Pro His Val Leu Ser Ser Glu Leu Asn Leu Ala Gly Ser 70 75 Ser Phe Val Val Cys Asp Val Ser Ser Leu Pro Gly Phe Ser Thr Ala 50 90 Lys Ser Glu Gly Ser Gly Val Thr Phe Leu Leu Gly Thr Lys Asp Ala 100 105 Glu Ala Ala Val Ala Lys Ala Val Asp Ala Gly Ala Val Lys Val Glu 120 125 55 Val Thr Glu Ala Glu Val Glu Leu Gly Phe Lys Gly Lys Val Thr Asp 140 135 Pro Phe Gly Val Thr Trp Ile Phe Ala Glu Lys Lys Thr Val Ile Thr

155

165
(2) INFORMATION FOR SEQ ID NO:143:
(i) SEQUENCE CHARACTERISTICS:

Asp Glu Asn Lys Glu Val

60

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(A) LENGTH: 138 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
 5
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..138
              (D) OTHER INFORMATION: / Ceres Seq. ID 2047216
10
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:143:
     Met Leu Leu Val Glu Ala Gln Lys Val Gly Asp Ala Val Thr Phe Tyr
                                      10
     Lys Ser Ala Phe Gly Ala Ile Glu Ser Gly His Ser Leu Tyr Pro Lys
                               25
15
     Arg Lys Leu Asp Gln Glu Leu Pro His Val Leu Ser Ser Glu Leu Asn
                               40
     Leu Ala Gly Ser Ser Phe Val Val Cys Asp Val Ser Ser Leu Pro Gly
     Phe Ser Thr Ala Lys Ser Glu Gly Ser Gly Val Thr Phe Leu Leu Gly
20
                                          7.5
     Thr Lys Asp Ala Glu Ala Ala Val Ala Lys Ala Val Asp Ala Gly Ala
                   85
                                      90 95
     Val Lys Val Glu Val Thr Glu Ala Glu Val Glu Leu Gly Phe Lys Gly
             100 105 110
25
     Lys Val Thr Asp Pro Phe Gly Val Thr Trp Ile Phe Ala Glu Lys Lys
          115 120 125
     Thr Val Ile Thr Asp Glu Asn Lys Glu Val
              135
        130
     (2) INFORMATION FOR SEQ ID NO:144:
30
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 98 amino acids
              (B) TYPE: amino acid
              (C) STRANDEDNESS:
              (D) TOPOLOGY: linear
35
         (ii) MOLECULE TYPE: peptide
         (ix) FEATURE:
              (A) NAME/KEY: peptide
              (B) LOCATION: 1..98
              (D) OTHER INFORMATION: / Ceres Seq. ID 2047217
40
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:144:
     Met Leu Leu Leu Leu Leu Thr Val Leu Val Arg Trp Arg Arg Ile
                                      10
     Leu Ser Ser Leu Ser Ser Arg Cys Cys Ser Leu Lys Leu Lys Arg
                                  25
45
     Ser Val Thr Leu Leu Ser Thr Asn Leu Leu Val Arg Ser Ser
     Leu Val Ile Leu Phe Thr Leu Ser Val Ser Leu Thr Lys Ser Phe Leu
     Met Phe Ser Leu Ser Leu Ile Ser Leu Ala Leu Pro Ser Leu Phe
50
     Ala Thr Phe Pro Leu Ser Leu Val Phe Leu Leu Arg Asn Arg Lys Val
     Arg Glu
55
     (2) INFORMATION FOR SEQ ID NO:145:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 682 base pairs
              (B) TYPE: nucleic acid
              (C) STRANDEDNESS: single
60
              (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: DNA (genomic)
```

(ix) FEATURE:

120

180 240

300

360

420

480

540

600

660

657

```
(A) NAME/KEY: -
                (B) LOCATION: 1..682
                (D) OTHER INFORMATION: / Ceres Seq. ID 2047438
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:145:
     atgattggtg gtttcttcgc cgatgtaaaa ttcaacagat cgagagagaa aaaactctct
     ctccctcctc cgttcaccgt cgcgactcgg ctctacagta cattttgttg agcaatctcg
     tatttagagg attccaatgt ctggcgcaga ggataataaa tcttcccatg ctcagctttc
     gtcccaaatt tttcttgatc ttgtggattc ggtgatagct gatgtagcat ctgagtgtca
      tcgggtagca cgactaggac ttgatcgtga tttggatata qtagaagaag agttgagatt
10
     qtcaqtqqaa qcqcqtqcqa aqatcqctqa tcctaqcaac aacctcqaaa ccaacaccaa
     atatgttgtt gatatatttg ggcagactca ccctcctgta gctagtgaag tgttcaattg
     catgaactgt qggcgacaaa ttgttgcagq aaqgtttgct cctcatctcg aaaaatgcat
      gggaaaggga agaaaggctc gtgccaaqac aaccaqaaqc acaacggctg cacagaaccg
      gaatqcacqa cqcaqcccca atccacqata ttctccttat ccaaattctq ctaqtqaqaa
15
      ccagttagca agtggatcac ctggtgttgc aggtgaagac tgctcaaatt tcacagttcg
      agagaacqtg aaaggagact ga
      (2) INFORMATION FOR SEQ ID NO:146:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 181 amino acids
20
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
25
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..181
                (D) OTHER INFORMATION: / Ceres Seq. ID 2047439
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:146:
     Met Ser Gly Ala Glu Asp Asn Lys Ser Ser His Ala Gln Leu Ser Ser
30
                                          10
      Gln Ile Phe Leu Asp Leu Val Asp Ser Val Ile Ala Asp Val Ala Ser
                                      25
      Glu Cys His Arg Val Ala Arg Leu Gly Leu Asp Arg Asp Leu Asp Ile
                                  40
35
      Val Glu Glu Glu Leu Arg Leu Ser Val Glu Ala Arg Ala Lys Ile Ala
                              55
      Asp Pro Ser Asn Asn Leu Glu Thr Asn Thr Lys Tyr Val Val Asp Ile
                          70
                                               75
      Phe Gly Gln Thr His Pro Pro Val Ala Ser Glu Val Phe Asn Cys Met
40
      Asn Cys Gly Arg Gln Ile Val Ala Gly Arg Phe Ala Pro His Leu Glu
                                      105
      Lys Cys Met Gly Lys Gly Arg Lys Ala Arg Ala Lys Thr Thr Arg Ser
                                  120
45
      Thr Thr Ala Ala Gln Asn Arg Asn Ala Arg Arg Ser Pro Asn Pro Arg
                              135
                                                   140
      Tyr Ser Pro Tyr Pro Asn Ser Ala Ser Glu Asn Gln Leu Ala Ser Gly
                          150
                                               155
      Ser Pro Gly Val Ala Gly Glu Asp Cys Ser Asn Phe Thr Val Arg Glu
50
                      165
                                          170
      Asn Val Lys Gly Asp
                  180
      (2) INFORMATION FOR SEQ ID NO:147:
           (i) SEQUENCE CHARACTERISTICS:
55
                (A) LENGTH: 501 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
60
          (ix) FEATURE:
                (A) NAME/KEY: -
```

(B) LOCATION: 1..501

```
658
                (D) OTHER INFORMATION: / Ceres Seq. ID 2049056
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:147:
      aaacaaaggt gtcaagaaga aggatcaaca tattaattga ctaaaaatga aggcatggat
      gataatcttg ttggtgattt gtgtcgctgt ggtggtggag caatcagagg ctcgcaaagg
                                                                             120
 5
      togaaagtat ttaaatocag gogtgottgamooggtgtogt ggtootaato otocagoggg
      atgtcatcct cacaattccc accacaaacc tcgcgtccct gttcacaatt atagtcgtgg
                                                                             240
      ttgtagtaga attacccggt gcagacgaga tgcctaggtt taacctcaca tgacgacgac
                                                                             300
                                                                             360
      gtccctttat attgatcctt cttctacatg agacttgcaa tgatattata tattcatata
      tatacacatt acqctqtata tqataattcc agttqattaa tatataaatc ttactqattt
                                                                             420
10
      caattcaata cacaagatcc tgtaactgaa ataattttgt ccatgttttg tgcttataat
                                                                             480
      aatggtatga ctcttattat t
      (2) INFORMATION FOR SEQ ID NO:148:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 76 amino acids
15
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
           ix) FEATURE:
20
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..76
                (D) OTHER INFORMATION: / Ceres Seq. ID 2049057
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:148:
      Met Lys Ala Trp Met Ile Ile Leu Leu Val Ile Cys Val Ala Val Val
25
                                          10
                     5
      Val Glu Gln Ser Glu Ala Arg Lys Gly Arg Lys Tyr Leu Asn Pro Gly
                                      25
                  20
      Val Leu Asp Arg Cys Arg Gly Pro Asn Pro Pro Ala Gly Cys His Pro
              35
                                  40
                                                       45
30
      His Asn Ser His His Lys Pro Arg Val Pro Val His Asn Tyr Ser Arg
                              55
      Gly Cys Ser Arg Ile Thr Ara Cys Arg Arg Asp Ala
                          7.0
      (2) INFORMATION FOR SEQ ID NO:149:
35
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 72 amino acids .
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
40
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..72
                (D) OTHER INFORMATION: / Ceres Seq. ID 2049058
45
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:149:
      Met Ile Ile Leu Leu Val Ile Cys Val Ala Val Val Glu Gln Ser
                                          10
                                                               15
      Glu Ala Arg Lys Gly Arg Lys Tyr Leu Asn Pro Gly Val Leu Asp Arg
                  20
                                       25
50
      Cys Arg Gly Pro Asn Pro Pro Ala Gly Cys His Pro His Asn Ser His
                                  40
      His Lys Pro Arg Val Pro Val His Asn Tyr Ser Arg Gly Cys Ser Arg
          50
                              55
                                                   60
      Ile Thr Arg Cys Arg Arg Asp Ala
55
                          70
      (2) INFORMATION FOR SEQ ID NO:150:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1007 base pairs
                (B) TYPE: nucleic acid
60
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
```

(ii) MOLECULE TYPE: DNA (genomic)

659

210

```
(ix) FEATURE:
                (A) NAME/KEY: -
                (B) LOCATION: 1..1007
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050386
 5
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:150:
     ccaaaatttct ccatggatcg ttctttgaat ctcctcgatt tagccttagg gttcgatgag
     cagctagcta ttccatcgcc actaaatgga aaagtaatac taatagaaga ctgtgtagag
                                                                            120
     acgagtggtt cctttgtact tcaccagcta atgaaacgtg ttctctcctc taactcctcc
                                                                            180
     gacgeastta tetttetege tittgetege cetttetete attatgateg aatettgegt
                                                                            240
10
     aaactgggat gtaatttagc tacccataag tcgaataatc gattggtgtt ctttgacatg
                                                                            300
     ctcatggtta agtgttcaga tggggatcaa atggaagaca atgtgagtgc agttgcgaaa
                                                                            360
     ctatttcggg agatacaaga aaccgttcga aagctacaga gtgtaacaag tggtaacata
                                                                            420
     actgttatgg tggatgacat gtctctgctg gaaattgcta ctaccggcag caactcagat
                                                                            480
     cacqtattgg acttettgca ttattgccac acattaaqtt ctgaaagcaa ttgttcattg
                                                                            540
15
                                                                            600
     gtcatcctca atcatgaaga tatatacqcg aqcatggaga gacctgcatt tttgctacag
     atggtatqcc ttqcaqatqt tqtqataaaq qcaqaqcctt taqcctctqq tttaqcaaat
                                                                            660
     gatgtacatg gccaattgac tgttctgaac aaagggataa gcaactcagg tagaggaagc
                                                                            720
     togaggaaca agttgcagaa ttttcaattc aggatcaaag aaaatggtat cgactatttc
                                                                            780
      tatcctqqtt qcaqaaqctq aqqattaqac accqttctqa qcattaqcat qqtaqtqaqa
20
      totttttcta gacagttttg ttatttgcgt ttaggttaaa cagagacact ctgttaaaca
     tttttacttt agtgtotott otggttgtaa ottgatgact tagotaaagt tgaatgottg
     agacacagaa acttgtagac atggaaaatt agtaacattc tatatct
      (2) INFORMATION FOR SEQ ID NO:151:
           (i) SEQUENCE CHARACTERISTICS:
25
                (A) LENGTH: 266 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
30
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..266
                (D) OTHER INFORMATION: / Ceres Seq. ID 2050387
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:151:
35
      Pro Asn Phe Ser Met Asp Arg Ser Leu Asn Leu Leu Asp Leu Ala Leu
                                          10
      Gly Phe Asp Glu Gln Leu Ala Ile Pro Ser Pro Leu Asn Gly Lys Val
                  20
                                      25
      Ile Leu Ile Glu Asp Cys Val Glu Thr Ser Gly Ser Phe Val Leu His
40
                                  40
      Gln Leu Met Lys Arg Val Leu Ser Ser Asn Ser Ser Asp Ala Leu Ile
                              55
      Phe Leu Ala Phe Ala Arg Pro Phe Ser His Tyr Asp Arg Ile Leu Arg
                          70
                                              75
45
      Lys Leu Gly Cys Asn Leu Ala Thr His Lys Ser Asn Asn Arg Leu Val
                                          90
      Phe Phe Asp Met Leu Met Val Lys Cys Ser Asp Gly Asp Gln Met Glu
                  100
                                      105
                                                          110
      Asp Asn Val Ser Ala Val Ala Lys Leu Phe Arg Glu Ile Gln Glu Thr
50
                                  120
                                                      125
      Val Arg Lys Leu Gln Ser Val Thr Ser Gly Asn Ile Thr Val Met Val
                              135
                                                  140
      Asp Asp Met Ser Leu Leu Glu Ile Ala Thr Thr Gly Ser Asn Ser Asp
                          150
                                              155
55
      His Val Leu Asp Phe Leu His Tyr Cys His Thr Leu Ser Ser Glu Ser
                                          170
                      165
      Asn Cys Ser Leu Val Ile Leu Asn His Glu Asp Ile Tyr Ala Ser Met
                                      185
      Glu Arg Pro Ala Phe Leu Leu Gln Met Val Cys Leu Ala Asp Val Val
60
                                                      205
                                 200
      Ile Lys Ala Glu Pro Leu Ala Ser Gly Leu Ala Asn Asp Val His Gly
```

215

660 Gln Leu Thr Val Leu Asn Lys Gly Ile Ser Asn Ser Gly Arg Gly Ser 230 235 Ser Arg Asn Lys Leu Gln Asn Phe Gln Phe Arg Ile Lys Glu Asn Gly 245 250 5 Ile Asp Tyr Phe Tyr Pro Gly Cys Arg Ser 260 265 (2) INFORMATION FOR SEQ ID NO:152: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 262 amino acids 10 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 15 (A) NAME/KEY: peptide (B) LOCATION: 1..262 (D) OTHER INFORMATION: / Ceres Seq. ID 2050388 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:152: Met Asp Arg Ser Leu Asn Leu Leu Asp Leu Ala Leu Gly Phe Asp Glu 20 1 5 10 Gln Leu Ala Ile Pro Ser Pro Leu Asn Gly Lys Val Ile Leu Ile Glu 25 Asp Cys Val Glu Thr Ser Gly Ser Phe Val Leu His Gln Leu Met Lys 40 25 Arg Val Leu Ser Ser Asn Ser Ser Asp Ala Leu Ile Phe Leu Ala Phe 55 Ala Arg Pro Phe Ser His Tyr Asp Arg Ile Leu Arg Lys Leu Gly Cys 70 Asn Leu Ala Thr His Lys Ser Asn Asn Arg Leu Val Phe Phe Asp Met 30 90 95 Leu Met Val Lys Cys Ser Asp Gly Asp Gln Met Glu Asp Asn Val Ser 105 Ala Val Ala Lys Leu Phe Arg Glu Ile Gln Glu Thr Val Arg Lys Leu 120 Gln Ser Val Thr Ser Gly Asn Ile Thr Val Met Val Asp Asp Met Ser 35 135 140 Leu Leu Glu Ile Ala Thr Thr Gly Ser Asn Ser Asp His Val Leu Asp 150 155 160 Phe Leu His Tyr Cys His Thr Leu Ser Ser Glu Ser Asn Cys Ser Leu 40 165 170 175 Val Ile Leu Asn His Glu Asp Ile Tyr Ala Ser Met Glu Arg Pro Ala 180 185 Phe Leu Leu Gln Met Val Cys Leu Ala Asp Val Val Ile Lys Ala Glu 200 45 Pro Leu Ala Ser Gly Leu Ala Asn Asp Val His Gly Gln Leu Thr Val 215 Leu Asn Lys Gly Ile Ser Asn Ser Gly Arg Gly Ser Ser Arg Asn Lys 230 235 Leu Gln Asn Phe Gln Phe Arg Ile Lys Glu Asn Gly Ile Asp Tyr Phe 50 245 250 Tyr Pro Gly Cys Arg Ser 260 (2) INFORMATION FOR SEQ ID NO:153: (i) SEQUENCE CHARACTERISTICS: 55 (A) LENGTH: 216 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 60 (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..216

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(D) OTHER INFORMATION: / Ceres Seq. ID 2050389
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:153:
     Met Lys Arg Val Leu Ser Ser Asn Ser Ser Asp Ala Leu Ile Phe Leu
                                          10
 5
     Ala Phe Ala Arg Pro Phe Ser His Tyr Asp Arg Ile Leu Arg Lys Leu
                                      25
     Gly Cys Asn Leu Ala Thr His Lys Ser Asn Asn Arg Leu Val Phe Phe
                                  40
      Asp Met Leu Met Val Lys Cys Ser Asp Gly Asp Gln Met Glu Asp Asn
10
                             55
      Val Ser Ala Val Ala Lys Leu Phe Arg Glu Ile Gln Glu Thr Val Arg
                         70
                                              75
      Lys Leu Gln Ser Val Thr Ser Gly Asn Ile Thr Val Met Val Asp Asp
                     8.5
                                          90
15
     Met Ser Leu Glu Ile Ala Thr Thr Gly Ser Asn Ser Asp His Val
                  100
                                      105
                                                          110
      Leu Asp Phe Leu His Tyr Cys His Thr Leu Ser Ser Glu Ser Asn Cys
                                  120
                                                      125
      Ser Leu Val Ile Leu Asn His Glu Asp Ile Tyr Ala Ser Met Glu Arg
20
                              135
      Pro Ala Phe Leu Gen Met Val Cys Leu Ala Asp Val Val Ile Lys
                          150
                                              155
      Ala Glu Pro Leu Ala Ser Gly Leu Ala Asn Asp Val His Gly Gln Leu
                      165
                                          170
25
      Thr Val Leu Asn Lys Gly Ile Ser Asn Ser Gly Arg Gly Ser Ser Arg
                                      185
      Asn Lys Leu Gln Asn Phe Gln Phe Arg Ile Lys Glu Asn Gly Ile Asp
              195
                                  200
      Tyr Phe Tyr Pro Gly Cys Arg Ser
30
                              215
      (2) INFORMATION FOR SEQ ID NO:154:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 634 base pairs
                (B) TYPE: nucleic acid
35
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: -
40
                (B) LOCATION: 1..634
                (D) OTHER INFORMATION: / Ceres Seq. ID 2053353
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:154:
      gaataaaaat ctatttcttg tgttttgtgtt gtatttactt aaaaaaataa aggagatcag
                                                                             60
      tttgggagaa agcagcaaag aagaaaaatg gcagcaacat ttgcaacacc atcgacggtg
                                                                             120
45
      ataggeeteg gaggateate cateaceace aaaccettet etteateett tttaaaacea
                                                                             180
      acattaagcg ccaagaaccc tttgagactc gccggtgcat cgggaggaag agtcacttgc
                                                                             240
      tttgagagga actggttgag gagagatttg aacgtggtag gatttgggct gatcggatgg
                                                                             300
      ctagctccgt cgagcattcc agcgataaat gggaagagcc tgacgggtct cttcttcgat
                                                                             360
      agcateggaa etgagetege teaetteeeg acteeteeag eteteaette acagttetgg
                                                                             420
50
      ttgtggttgg ttacgtggca cttaggcctc ttcctctgcc tcactttcgg acaaatcgga
                                                                             480
      ttcaagggca ggactgaaga ttacttctaa ggataactat tcttgttttc gtttgtacta
                                                                             540
      tatgctctct cttggttatg tgtaatatta tcaatcaaaa caaagctttc tttgctattt
                                                                             600
      gatgtttcat tctatatctc gacaagtctt tctt
      (2) INFORMATION FOR SEQ ID NO:155:
55
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 140 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
60
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
```

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(B) LOCATION: 1..140
                (D) OTHER INFORMATION: / Ceres Seq. ID 2053354
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:155:
     Met Ala Ala Thr Phe Ala Thr Pro Ser Thr Val Ile Gly Leu Gly Gly
 5
                     5
                                        10
      Ser Ser Ile Thr Thr Lys Pro Phe Ser Ser Ser Phe Leu Lys Pro Thr
                                      25
      Leu Ser Ala Lvs Asn Pro Leu Arg Leu Ala Gly Ala Ser Gly Gly Arg
                                  40
10
      Val Thr Cys Phe Glu Arg Asn Trp Leu Arg Arg Asp Leu Asn Val Val
                              55
      Gly Phe Gly Leu Ile Gly Trp Leu Ala Pro Ser Ser Ile Pro Ala Ile
                          70
                                              75
      Asn Gly Lys Ser Leu Thr Gly Leu Phe Phe Asp Ser Ile Gly Thr Glu
15
                      8.5
                                          90
      Leu Ala His Phe Pro Thr Pro Pro Ala Leu Thr Ser Gln Phe Trp Leu
                  100
                                      105
      Trp Leu Val Thr Trp His Leu Gly Leu Phe Leu Cys Leu Thr Phe Gly
                                  120 125 .
20
      Gln Ile Gly Phe Lys Gly Arg Thr Glu Asp Tyr Phe
          130
                              135
      (2) INFORMATION FOR SEQ ID NO:156:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 393 base pairs
25
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
30
                (A) NAME/KEY: -
                (B) LOCATION: 1..393
                (D) OTHER INFORMATION: / Ceres Seq. ID 2055693
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:156:
      atgcaaaaac ctctctctct ctctcaattc acttctcctc cttcaatggc ttggcgcaac
                                                                             60
35
      gcaggatetg ecgetegtte tttegtetee gceaeegeaa gateaecate teteegttet
                                                                            120
      cccaccacgg cgcttcctcg cctccgtcct cctcaatcct ccttacctag ccgtcgcttc
                                                                            180
                                                                            240
      acctteteat cacctteeag gaatetagga geacttggtt geacacagte gttettgeet
                                                                            300
      ctgtacagtg ttgtggctac ttctcaactc acatctcacc ttaacgttaa cttgcgagct
                                                                            360
      ttotgogago tgtotaacgg gaattggaaa agatgggtga tgcactogga ttttacgtog
40
      ctggcaagag aatggtgctg cattgttccg tga
      (2) INFORMATION FOR SEQ ID NO:157:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 130 amino acids
                (B) TYPE: amino acid
45
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
50
                (B) LOCATION: 1..130
                (D) OTHER INFORMATION: / Ceres Seq. ID 2055694
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:157:
      Met Gln Lys Pro Leu Ser Leu Ser Gln Phe Thr Ser Pro Pro Ser Met
                                          10
55
      Ala Trp Arg Asn Ala Gly Ser Ala Ala Arg Ser Phe Val Ser Ala Thr
                                      25
      Ala Arg Ser Pro Ser Leu Arg Ser Pro Thr Thr Ala Leu Pro Arg Leu
                                  40
      Arg Pro Pro Gln Ser Ser Leu Pro Ser Arg Arg Phe Thr Phe Ser Ser
60
                              55
                                                  60
      Pro Ser Arg Asn Leu Gly Ala Leu Gly Cys Thr Gln Ser Phe Leu Pro
                                              75
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663 Leu Tyr Ser Val Val Ala Thr Ser Gln Leu Thr Ser His Leu Asn Val 90 8.5 Asn Leu Arg Ala Phe Cys Glu Leu Ser Asn Gly Asn Trp Lys Arg Trp 105 100 Val Met His Ser Asp Phe Thr Ser Leu Ala Arg Glu Trp Cys Cys Ile 120 125 115 Val Pro 130 (2) INFORMATION FOR SEQ ID NO:158: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 115 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: peptide (ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..115 (D) OTHER INFORMATION: / Ceres Seq. ID 2055695 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:158: Met Ala Trp Arg Asn Ala Gly Ser Ala Ala Arg Ser Phe Val Ser Ala 5 10 Thr Ala Arg Ser Pro Ser Leu Arg Ser Pro Thr Thr Ala Leu Pro Arg 20 25 25 Leu Arg Pro Pro Gln Ser Ser Leu Pro Ser Arg Arg Phe Thr Phe Ser 40 Ser Pro Ser Arg Asn Leu Gly Ala Leu Gly Cys Thr Gln Ser Phe Leu 55 Pro Leu Tyr Ser Val Val Ala Thr Ser Gln Leu Thr Ser His Leu Asn 30 Val Asn Leu Arg Ala Phe Cys Glu Leu Ser Asn Gly Asn Trp Lys Arg 90 Trp Val Met His Ser Asp Phe Thr Ser Leu Ala Arg Glu Trp Cys Cys 105 35 Ile Val Pro 115 (2) INFORMATION FOR SEQ ID NO:159: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 68 amino acids 40 (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (ix) FEATURE: 45 (A) NAME/KEY: peptide (B) LOCATION: 1..68 (D) OTHER INFORMATION: / Ceres Seq. ID 2055696 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:159: Cys Lys Asn Leu Ser Leu Ser Leu Asn Ser Leu Leu Leu Gln Trp 50 10 Leu Gly Ala Thr Gln Asp Leu Pro Leu Val Leu Ser Ser Pro Pro 20 25 Gln Asp His His Leu Ser Val Leu Pro Pro Arg Arg Phe Leu Ala Ser 40 45 55 Val Leu Leu Asn Pro Pro Tyr Leu Ala Val Ala Ser Pro Ser His His 55 Leu Pro Gly Ile (2) INFORMATION FOR SEQ ID NO:160: 60 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 394 base pairs (B) TYPE: nucleic acid

WO 00/40095

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664
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
 5
                (A) NAME/KEY: -
                (B) LOCATION: 1..394
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056405
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:160:
     tcaatagett atctagttet tgactgttea acgateaatg geaccateeg etgeaatget
                                                                             60
10
     catactetea cateetetag ttagecacaa agetaagaat cagtetetgt categeegte
                                                                             120
      gtctgttaag tcgacacgtg tcttcggttt tctttggcct tggaaggcat tagacaatga
                                                                             180
     ggatcattcc gcggttgttc tcggccggct ctttgqcqat ccggctacta tcgagaagcg
                                                                             240
      cttccaagaa gctcttgaac aaagctgttg gtaatttgta caatgtttgg ttgggtttat
                                                                             300
      tgattgtttt tacgttaaaa tagcttttat aaattggaaa tggaagtact gtaaaatgta
15
      aaaattgact atatataatt aaaggtacat tatg
      (2) INFORMATION FOR SEQ ID NO:161:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 78 amino acids
                (B) TYPE: amino acid
20
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                (A) NAME/KEY: peptide
25
                (B) LOCATION: 1..78
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056406
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:161:
      Met Ala Pro Ser Ala Ala Met Leu Ile Leu Ser His Pro Leu Val Ser
                      5
                                          10
                                                               15
30
      His Lys Ala Lys Asn Gln Ser Leu Ser Ser Pro Ser Ser Val Lys Ser
                  20
                                      25
      Thr Arg Val Phe Gly Phe Leu Trp P:o Trp Lys Ala Leu Asp Asn Glu
                                  40
      Asp His Ser Ala Val Val Leu Gly Arg Leu Phe Gly Asp Pro Ala Thr
35
      Ile Glu Lys Arg Phe Gln Glu Ala Leu Glu Gln Ser Cys Trp
                          70
      (2) INFORMATION FOR SEO ID NO:162:
           (i) SEQUENCE CHARACTERISTICS:
40
                (A) LENGTH: 72 amino acids
                (B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
45
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
                 (B) LOCATION: 1..72
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056407
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:162:
50
      Met Leu Ile Leu Ser His Pro Leu Val Ser His Lys Ala Lys Asn Gln
                                          10
                     5
      Ser Leu Ser Ser Pro Ser Ser Val Lys Ser Thr Arg Val Phe Gly Phe
                  20
                                      25
      Leu Trp Pro Trp Lys Ala Leu Asp Asn Glu Asp His Ser Ala Val Val
55
                                  40
      Leu Gly Arg Leu Phe Gly Asp Pro Ala Thr Ile Glu Lys Arg Phe Gln
                              55
      Glu Ala Leu Glu Gln Ser Cys Trp
                          70
60
      (2) INFORMATION FOR SEQ ID NO:163:
           (i) SEQUENCE CHARACTERISTICS:
```

(A) LENGTH: 48 amino acids

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(B) TYPE: amino acid
                (C) STRANDEDNESS:
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
 5
          (ix) FEATURE:
                (A) NAME/KEY: peptide
                (B) LOCATION: 1..48
                (D) OTHER INFORMATION: / Ceres Seq. ID 2056408
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:163:
10
     Met Arg Ile Ile Pro Arg Leu Phe Ser Ala Gly Ser Leu Ala Ile Arg
                      5
                                          10
      Leu Leu Ser Arg Ser Ala Ser Lys Lys Leu Leu Asn Lys Ala Val Gly
                                      25
      Asn Leu Tyr Asn Val Trp Leu Gly Leu Leu Ile Val Phe Thr Leu Lys
15
              35
                                  40
      (2) INFORMATION FOR SEQ ID NO:164:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 766 base pairs
                (B) TYPE: nucleic acid
20
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: DNA (genomic)
          (ix) FEATURE:
                (A) NAME/KEY: ~
25
                (B) LOCATION: 1..766
                (D) OTHER INFORMATION: / Ceres Seq. ID 2065747
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:164:
      acagtegeet cetteetgeg eegeegeege egeegeeget accaeegeet egeggeegee
                                                                              60
      tgcggctccg cgtttccgca cggatttggt gaaggtttct tgaggctgag aacatggtgc
                                                                             120
30
      tgaagacgga actttgccgc ttcagcggcc agaagattta tcctgggaaa ggcattagat
                                                                             180
                                                                             240
      ttatccgtgc tgattctcag gtcttccttt ttgccaactc gaaatgcaag cgctacttcc
                                                                             300
      acaaccgcct gaagcctgca aagcttacct çgacagcaat gtacaggaag cagcacaaga
                                                                             360
      aggatatcca tgctgaagcg gtaaagaaga ggcgccgcgc caccaagaag ccatactcca
      ggtcaattgt gggtgcttcc ttggaagtaa tccagaagaa gagagctgag aagccagagg
                                                                             420
35
      tecgegatge tgetagagaa getgetette gtgagateaa ggagegeate aagaagaeea
                                                                             480
                                                                             540
      aggatgagaa gaaagcgaag aaggcggagg tgagcaagtc ccagaagacg cagacaaagg
                                                                             600
      gtgcggtcca gaagggttcc aagggcccca agttgggcgg cggtggtggg aagcgctgaa
      agaacttagt gtcgtttctc gacattgcag tcgttcctta gccaaagcca ctttcgtaga
                                                                             660
                                                                             720
      acctgtgttg aatttgcaag acttattcaa gcgttgcttg tgcgtgctaa ataccatggc
40
      aagagaacgg atttatattt atgcctgaaa aaaaatgacc gttcat
      (2) INFORMATION FOR SEQ ID NO:165:
           (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 161 amino acids
                 (B) TYPE: amino acid
45
                 (C) STRANDEDNESS:
                 (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
          (ix) FEATURE:
                 (A) NAME/KEY: peptide
50
                 (B) LOCATION: 1..161
                 (D) OTHER INFORMATION: / Ceres Seq. ID 2065748
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:165:
      Met Val Leu Lys Thr Glu Leu Cys Arg Phe Ser Gly Gln Lys Ile Tyr
                                           10
55
      Pro Gly Lys Gly Ile Arg Phe Ile Arg Ala Asp Ser Gln Val Phe Leu
                                       25
      Phe Ala Asn Ser Lys Cys Lys Arg Tyr Phe His Asn Arg Leu Lys Pro
                                   40
      Ala Lys Leu Thr Trp Thr Ala Met Tyr Arg Lys Gln His Lys Lys Asp
60
                               55
                                                   60
      Ile His Ala Glu Ala Val Lys Lys Arg Arg Ala Thr Lys Lys Pro
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666 Tyr Ser Arg Ser Ile Val Gly Ala Ser Leu Glu Val Ile Gln Lys Lys 90 Arg Ala Glu Lys Pro Glu Val Arg Asp Ala Ala Arg Glu Ala Ala Leu 100 105 110 Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala 115 120 125 Lys Lys Ala Glu Val Ser Lys Ser Gln Lys Thr Gln Thr Lys Gly Ala 130 135 140 Val Gln Lys Gly Ser Lys Gly Pro Lys Leu Gly Gly Gly Gly Lys 10 155 150 Arq (2) INFORMATION FOR SEQ ID NO:166: (i) SEQUENCE CHARACTERISTICS: 15 (A) LENGTH: 106 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 20 ix) FEATURE: (A) NAME/KEY: peptide (B) LOCATION: 1..106 (D) OTHER INFORMATION: / Ceres Seq. ID 2065749 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:166: 25 Met Tyr Arg Lys Gln His Lys Lys Asp Ile His Ala Glu Ala Val Lys 10 5 Lys Arg Arg Arg Ala Thr Lys Lys Pro Tyr Ser Arg Ser Ile Val Gly 20 25 Ala Ser Leu Glu Val Ile Gln Lys Lys Arg Ala Glu Lys Pro Glu Val 30 40 Arg Asp Ala Ala Arg Glu Ala Ala Leu Arg Glu Ile Lys Glu Arg Ile Lys Lys Thr Lys Asp Glu Lys Lys Ala Lys Lys Ala Glu Val Ser Lys 75 Ser Gln Lys Thr Gln Thr Lys Gly Ala Val Gln Lys Gly Ser Lys Gly 35 85 Pro Lys Leu Gly Gly Gly Gly Lys Arg 40

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CLAIMS

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What is claimed is:

1. An isolated nucleic acid molecule comprising a nucleic acid having a nucleotide sequence selected from the group consisting of

- (i) a nucleotide sequence shown in SEQ TABLES 1 AND 2;
- (ii) a nucleotide sequence complementary to one shown in SEQ TABLES 1 AND 2;
  - (iii) a nucleotide sequence which is the reverse complement of one shown in SEQ TABLES 1 AND 2;
- (iv) a nucleotide sequence capable of hybridizing to a 10 nucleotide sequence shown in SEQ TABLES 1 AND 2;
  - (iv) a nucleotide sequence capable of hybridizing to a nucleotide sequence complementary to one shown in SEQ TABLES 1 AND 2;
- (vi) a nucleotide sequence capable of hybridizing to a
  15 nucleotide sequence which is the reverse complement of one
  shown in SEQ TABLES 1 AND 2;

whereby the hybridization in groups (iv) to (vi) allow said sequences to form a duplex at a temperature from about  $Tm-40\,^{\circ}C$  to about  $Tm-48\,^{\circ}$  C.

- 2. An isolated nucleic acid molecule according to claim 1 that comprises a complete open reading frame.
- 3. An isolated nucleic acid molecule according to claim 1 that functions as a promoter or as a 3' end termination sequence or as a regulator sequence influencing the transcription rate, the transcript stability or RNA translation rate in a host cell.

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4. The isolated nucleic acid of claim 3, comprising a sequence selected from the group consisting of a TATA box sequence, a CAAT box sequence, a motif of GCAATCG or a transcription-factor binding sequence.

5.An isolated nucleic acid molecule comprising a nucleic acid sequence that encodes an amino acid sequence selected from the group consisting of the amino acid sequences shown in SEQ TABLES 1 AND 2, a complementary nucleotide sequence to said nucleic acid sequence that encodes said amino acid sequence selected from SEQ TABLES 1 AND 2, a reverse complementary nucleotide sequence to said nucleic acid sequence that encodes said amino acid sequence selected from SEQ TABLES 1 AND 2, and a nucleotide sequence that will hybridize to said nucleic acid sequence encoding an amino acid selected from SEQ TABLES 1 AND 2 or said complementary sequence under hybridization conditions 40°C Tm 48°C. providing Τm to

10

10

15

6. isolated nucleic acid molecule comprising An nucleic acid sequence that encodes at least a part or a portion or a mutant or a fusion of an amino acid sequence selected from the group consisting of the amino acid sequences shown in SEQ TABLES 1 AND 2, a complementary nucleotide sequence to said nucleic acid sequence that encodes said part or portion or mutant or fusion of an amino acid sequence selected from SEQ TABLES 1 AND 2, a reverse complementary nucleotide sequence to said nucleic acid sequence that encodes said part or portion or mutant or fusion of an amino acid sequence selected from SEQ TABLES 1 AND 2, and a nucleotide sequence that will hybridize to said nucleic acid sequence that encodes said part or portion or mutant or fusion of an amino acid selected from SEQ TABLES or said complementary sequence or said reverse complementary sequence under hybridization conditions providing Tm - 40°C to Tm - 48°C.

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- 7. An isolated nucleic acid molecule according to claim 1, having a nucleotide sequence selected from the group consisting of the nucleotide sequences shown in SEQ TABLES 1 AND 2, and a complementary nucleotide sequence to said nucleotide sequence selected from SEQ TABLES 1 AND 2.
- 8. A chimeric DNA construct comprising (a) a promoter sequence capable of driving gene expression in plant cells and operatively linked to (b) a structural gene comprising an DNA molecule according to any one of claims 1, 2, 5 and 7.
- 9. A chimeric DNA construct comprising (a) a promoter that is a nucleic acid molecule according to claim 3 or 4 operatively linked to (b) a structural gene and, optionally, (c) an associated UTR.
- 10. An isolated DNA molecule comprising (a) a promoter sequence according to claim 3 or 4 and operatively linked to (b) a structural gene sequence according to any one of claims 1, 2, 5, 6 and 7.
- 11. A promoter according to claim 10, wherein said promoter is a specific promoter.
- 12. A promoter according to claim 11, wherein said promoter is a seed-specific promoter, an embryo-specific promoter, an ovule-specific promoter, a tapetum-specific promoter or a root-specific promoter.
- 13. A host cell transformed with a polynucleotide comprising the isolated nucleic acid molecule according to claim 1.

- 14. An isolated polypeptide comprising an amino acid sequence encoded by a polynucleotide sequence shown in SEQ TABLES 1 AND 2, or one that is at least 75% identical thereto.
- 15. An isolated polypeptide according to claim 14, that is at least 85% identical to said amino acid sequence.
- 16. An isolated polypeptide, according to claim 15, that is at least 90% identical to said amino acid sequence.
- 17. A polynucleotide comprising a first polynucleotide sequence from SEQ TABLES 1 AND 2 or a fragment thereof, wherein said\_first sequence is capable of regulating transcription or translation of second a polynucleotide comprising a coding sequence in a host cell or *in vitro* system wherein the first and second polynucleotides are operatively linked.
- 18. The first polynucleotide of claim 17, which is capable of regulating transcription or translation in an *in vitro* system.
- 19. The first polynucleotide of claim 17, which is capable of regulating transcription or translation in a host cell.
- 20. The polynucleotide of claim 19, wherein said host cell is a plant cell.
- 21. A host cell of claim 13, wherein said isolated nucleic acid comprises a promoter and operatively linked structural gene and further wherein said promoter and structural gene are not heterologous to each other and are exogenous to the host cell genome.

- 22. A method of introducing an isolated nucleic acid into a host cell comprising:
  - (a) providing an isolated nucleic acid of any of claims 1-12:
- 5 (b) contacting said isolated nucleic acid with said host cell under conditions that permit insertion of said nucleic acid into said host cell.
  - 23. A method of producing a polypeptide of any one of claims 14-16 comprising:
    - (a) providing a host cell of claim 13;
- (b) culturing said host cell under conditions that permit transcription and translation of said structural gene to produce a polypeptide; and
  - (c) isolating said polypeptide.
  - 24. An antibody capable of binding to a polypeptide of any one of claims 14-16.
  - 25. An isolated nucleic acid comprising a promoter of a gene wherein said gene comprises a nucleic acid having the nucleotide sequence of claim 1.
  - 26. An isolated polypeptide comprising an amino acid sequence selected from SEQ TABLES 1 AND 2, or one that is at least 75% identical thereto.
  - 27. The isolated polypeptide according to Claim 26 that is at least 85% identical thereto.
  - 28. The isolated polypeptide according to claim 26 that is at least 90% identical thereto.